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and a light lastness stabilizer, and a process for preparing the decolorizable toner, which comprises the steps of mixing the near intrared ray - absorbing dye and the light fashness stabilizer with the resin binder, and then mixing the decolorizing agent therewith. The decolorizable toner has excellent light stability during its production A decolorizable toner comprising a resin binder, a near intrared ray - absorbing dye, a decolorizing agent S Decolorizable toner.

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or storage and excellent decolorizing property.

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The present invention relates to a decolorizable toner, and more particularly to a near infrared ray decolorizable toner which can make an electric latent image or an electric signal used in electrophotog raphy, electrostatic recording materials and the like visible.

In recent years the reuse or recyle of used papers has been studied in order that lorest resources can be protected and that used papers can be diminished from cities. As a part of the reuse or recyle of used papers, the reuse of used papers such as copied paper, printed paper and used facsimile paper which are disused in an office of enterprise, has been studied.

fhere is an attempt to join a paper – manufacturing company in one enterprise so that used papors can be reused or recycled by collecting and dissolving the used papers to produce regenerated paper in the paper - manufacturing company However, since most of these used papers are generally confidential a paper - manufacturing company outside the enterprise. Also, since recorded figures or printed images papers of the enterprise, it is very difficult that the used papers are collected to produce regenerated paper Therefore, it has been thought that the used papers cannot be probably actually reused. Also, although the the used papers cannot be easity erased, the used papers are obliged to burn up or crush for dumping. recycle of the used papers which are previously crushed by using a paper-shredder has been studied. there is a defect that regenerated paper made of the crushed papers cannot be used as, for instance, specially papers for communication since the regenerated paper generally has small mechanical strength.

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As a toner which dissolves the above problems, we have developed a decolorizable toner containing a near infrared ray - decolorizing recording material which is decomposed and decolorized by absorbing near infrared rays.

When an electrostatic copying is carried out with the decolorizable toner, images or printed figures recorded on a recording paper can be easily decolorized only by the irradiation of near infrared rays, and therefore, the recording paper can be reused. Also, when the used recording paper is disposed, the recorded images or printed figures can be decolorized by the irradiation of near infrared rays, and therefore, there are many advantages such that the collected used recording papers can be reused at the same time that the transpiration of socret matters described in confidential papers can be prevented.

decolorizing material when the decolorizable toner is exposed to natural light during the production or storage of the decolorizable toner since the natural light contains a light which denaturalizes the near infrared ray -decolorizing material and a sensitizer for decolorizing the near infrared ray -decolorizing The decolorizable toner, however, has disadvantages in terms of decolorizing of the near infrared ray – material contained in the above decolorizable toner.

An object of the present invention is to provide a decolorizable toner having excellent light stability during the production or storage of the decolorizable toner, and forming images having excellent light resistance.

A further object of the present invention is to provide a decolorizable toner having excellent decolorizing property

These and other objects of the present invention will become apparent from the description hereinafter. In accordance with the present invention, there are provided

(1) a decolorizable toner comprising

(A) a resin binder.

(B) at least one near infrared ray - absorbing dye selected from the group consisting of a near infrared

ray - absorbing dye represented by the general formula (f);

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wherein X⁻ is a halogen ion perchloric acid ion, PF₆⁻, SbF₆⁻, OH⁻, sulfonic add ion or BF₆⁻, Y⁺ is a cation having absorptions in the near infrared region, and a near infrared ray absorbing dye epresented by the general formula (II)

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wherein each of H', H', H' and H' is independently hydrogen atom a hydrocarbon group or a hydrocarbon group containing a hotero atom, Y' is the same as defined above,

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(C) a decolorizing agent represented by the general formula (III):

 $\overline{\mathbb{B}}$

wherein each of R⁵. R⁶. R⁷ and R⁸ is independently an alkyl group, an aryl group, an allyl group, an aralkyl group, an alkenyl group, an alkynyl group, silyl group, an alicyclic group, a substituted alkyl group, a substituted aryl group, a substituted allyl group, a substituted aralkyl group, a substituted alkenyl group, a substituted alkynyl group or a substituted silyl group, with the proviso that at least one of R', R', R' and R' is an alkyl group having 1 to 12 carbon atoms; and each of R', R'', R'' and R^{12} is independently hydrogen atom, an atkyl group, an aryl group, an atlyl group, an aralkyl group, an alkenyl group, an alkynyl grout), an alicyclic group, a substituted alkyl group, a substituted aryl group, a substituted allyl group, a substituted aralkyl group, a substituted alkenyl group or a substituted

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(D) a light fastness stabilizer, and

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(2) a process for preparing a decotorizable toner which comprises the steps of mixing a near infrared ray - absorbing dye and a light tastness stabilizer with a resin binder, and then mixing a decolorizing

The decolorizable toner of the present invention is hardly discolored due to the denaturalization of a near infrared ray - absorbing dye contained in the decolorizable toner when the decolorizable toner is exposed to natural light during its production or storage. Therefore, the decolorizable toner shows excellent storage stability. Moreover, the images formed from the decolorizable toner have excellent light resistance 23

instance, polystyrane resins represented by polystyrene and the like, polyester resins represented by a Typical examples of the resin binder used in the decolorizable toner of the present invention are, for saturaled polyester, an unsaturated nolyester and the like: epoxy resins; (meth)acrylic resins represented by polymethyl methacrylate and the like; silicone resins; fluorocarbon resins; polyamide resins; polywinyl ins; rosin - denatured phenol - formaldehyde resins, and the like, and the present invention is not limited to alcohol resins; polyurethane resins; polyoletinic resins; polyvinylbulyrał resins; phenol - formaldehyde res the exemplified ones. These resin birders can be generally used alone or in admixture thereof. 8

When a resin having a melt viscosity of at most 10' poise at a temperature of 110°C and a melt toner, a printed image and the like, which are formed from the decolorizable toner on a support such as viscosity of all least 102 poise at a lemperature of 140°C is used as the resin binder of the decolorizable copying paper can be almost completely decolorized by irradiating near infrared rays. Also, after decolorizing. The decolorized printed image is hardly discolored when a visible light is irradiated to the decolorized printed image.

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When the printed image formed from the decolorizable toner is decolorized by means of, for instance, a halogen lamp and the like, the decolorizable toner is generally sometimes heated to 110°C or more. If the interaction of the near infrared ray-absorbing dye and the decolorizing agent is insufficient at that lemperature, decolorizing property is not sufficiently imparted to the decolorizable toner. When a resin binder having a specific viscosity at 110°C is used, since the interaction of the near infrared ray - absorbing dye and the decolorizing agent is maintained by the mobility of the resin binder, the decolorizing property is ş

The near infrared ray - absorbing dye is generally thermally decomposed at a temperature of around 140°C and decomposition products generated therefrom deteriorate visible light resistance of the de-

When a resin binder shows a sufliciently high melt viscosity at that temperature, since the near infrared ray -absorbing dye would not move in the tesin binder, the near infrared ray -absorbing dye is hardly decomposed by the heat.

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using the toner. For instance, when the resin binder has a melt viscosity of less than 10° poise at a Furthermore, if the mobility of the resin binder is remarkably changed by the change of a heating lemperature of the resul binder, a problem occurs in forming a figure or an image on a recording paper by lemperature of 140°C, the resin binder is completely molten by the heat for fixing the toner on a recording paper, and the interaction between the near infrared ray -absorbing dye and the decolorizing agent is promoted. As a result, the recorded figure or image is discolored when a visible light is irradiated to the 55

ligure or image. However, when a resin binder having a melt viscosity of al least 103 poise at a temperature of 140°C is used, the abuve phenoinenon does not occur and the docolorizable toner shows excellent

As mentioned above, in the present invention, it is preferred that a resin having a melt viscosity of al most 10' poise at a temperature of 110°C and a melt viscosity of at least 10' poise at a temperature of

The above - mentioned melt viscosity is intended to refer to a value measured by using a Shimadzu Flowlester. More particularly, the above - mentioned melt viscosily is intended to refer to a viscosity which is determined by charging a cylinder quipped with a die having a length of 1 mm and a diameter of 1 mm of Shirnadzu Flowtester CFT - 500 commercially available from SHIMADZU CORP, with 1.5 g of a resin binder, increasing the temperature of the resin binder at an increasing temperature of 2°C:minute while applying a pressure of 10 kg-t/cm. To the binder resin with a plunger, and measuring the viscosity of the resin binder at a temperature of 110°C or 140°C.

When the melt viscosity of the resin bunder is higher than 10' poise at 110°C, even if the temperature of the toner attains to around 110°C by the heat from a halogen lamp or the like during the decolorization absorbing dye and the decolorizing agent is lowered, and the improvement decolorizing property cannot be the figure or image formed from the toner is promoted by the irradiation of visible light. Accordingly, in the present invention, it is desired that a resin having a melt viscosity of at most 10' poise, preferably at most expected. When the melt viscosity of the resin binder is lower than 103 poise at 140°C, as mentioned above, the resin binder of the toner is excessively molten during fixing the toner, and the discolorization of the toner, the mobility of the resin binder is small. the contact frequency of the near infrared ray poise at a temperature of 110°C and a melt viscosity of at least 10³ poise, preferably at least $5 imes 10^3$ poise at a temperature of 140°C is used as the resin binder. 2

Concrete examples of the resin binder having the above-mentioned melt viscosity are, for instance, homopolymers of styrene or substituted styrenes such as polystyrene poly - p - chlorostyrene and poly vinylloluene: styrene copolymers such as a styrene - p - chlorostyrene copolymer, a styrene - propylene copolymer, a styrene - vinyttoluene copolymer, a styrene - vinylnaphthalene copolymer, a styrene - methyl octyl acrylate copolymer, a styrene-methyl methacrylate copolymer, a styrene-ethyl methacrylate isoprene copolymer, a styrene - acrylonitrile - indene copolymer, a styrene - maleic acid copolymer and a acrylate copolymer, a styrene – ethyl acrylate copolymer, a styrene – butyl acrylate copolymer, a styrene – copolymer, a styrene - butyl methacrylate copolymer, a styrene - methyl o - chloromethacrylate copolymer, a Slyrane - acrylonitrile copolymer, a styrene - vinyl methyl ether copolymer, a slyrene - vinyl ethyl ether styrene – maleic acid ester copolymer; potymethyl methacrylate, polybutyt methacrylate, polyvinyl chloride, invention is not limited to the exemplified ones. These, resin binders are generally used alone or in copolymer, a styrene-vnyl methyl ketone copolymer, a styrene-butadiene copolymer, a styrenepolyvinyl acetate, polyethylene, polypropylene, polyester, polywethane, polyamide, an epoxy resin, poly -vnył butyrał, an acrylic acıd resin, rosin, denatured rosin, a terpene resin, a phenol resin, an aliphatic or alicyclic hydrocarbon resin, an aromatic petroleum resin, chlorinaled paraffin, and the tike and the present admixture thereof. Among these resin binders, styrene - acrylic acid ester copolymers and polyester can be particularly preferably used since the near infrared ray -absorbing dye and the decoloring agent can be uniformly dispersed in these resins and therefore a decolorizable toner can be easily prepared therefrom, and further the decolorizable toner has excellent decolorizing property and discoloration resistance against £ 2 š ŝ

If is preferred that a thermoplastic resin having a light transmittance of at least 80 %, particularly at loast 85 % is used as a resin binder in order that affermages are scarcely remained after the decolorization ç

In the present specification, the light transmittance is intended to refer to a value measured by a method prescribed in JtS K 6717 Item 4.5 (1977).

In the present invention, the light transmittance was measured by using a direct reading haze meter Commercially available from TOYO SEIKI SEISAKU - SYO, LTD 7

Concrete examples of the thermoplastic resin are. for instance, polystyrene resins such as styrene

hydrogenated polystyrene, a styrene-propylene copolymer, a styrene-isobutylene Copolymer, a styrene - butadiene copolymer, a styrene - allyt alcohol copolymer, a styrene - maleic acid acrylic rubber-styrana torpolymer and an acrylonitrile-chlorinated polyethylene-styrene terpolymer; acrylic resins such as polymethyl methacrylate, polyethyl methacrylate, polybutyl methacrylate, polyglycidyl methacrylate, a lluorine - containing acrylate - methyl methacrylate - bulyl methacrylate copolymer and an an acrylonitiile - styrene - acrylic acid ester terpolymer, a styrene - acrylonitiile copolymer, an acrylonitiile osfer copolymer, a slyiene - maleic anhydride copolymer, an acrylonitrile - butadiene - styrene lerpolymer, homoplyrner,

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acrylic acid copolymer, a styrene-buladiene-acrylic acid ustiri, соробутег, а styrene-metry. methacrylate copolymer, a styrene-methyl methacrylate-bulyl acrylate terpolymor, a styrene-glycarlyr acrylic acid ester - maleic acid ester lerpolymer and a styrene - butyl acrylate - acrylic acid terpolymer, and methacrylate copolymer, a styrene - butadiene - dunethylaminoethyl muthacrylate terpolymer, a styrining othył aczylate - aczylic acid copolymer, styrone - (meth)aczylic acid uster copolymors such as a styrom Copolymer, a styrene-butyl muthacrylate copolymer, the like. These resins can be used alone or in admixture therior.

As the resin binder, a resin binder having a large polarity can be particularly proferably used since өжсөllөлт discoloration resistance is imparted to a toner. Examples of such resin binder are, for instan.e. resins having at least one group selected from hydroxyl group, cyano group, carboxyl group, carbuniyl group and ketone group in its molecule, such as polyester resins, epoxy resins, (meth)acryhic resins, polyamide resins, polyvinyl alcohol resins, polyurelhane resins, polyacrylonitrile resins, polyvinyl acetate resins. phenol resins. styrene - acrylic acid copolymers. styrene - acrylointrile copolymers, ethylone - viriyl acetate copolymers or ethylene - acrylic acid copolymers 5 2

The amount of the resin binder having a large potenty cannot be absolutely determined since the degree of potarity changes depending upon the kind of the polar group existing in the resin binder, but it is generally preferable that the content of such resin binder in the total amounts of the resin binder is at least 5 % by weight, particularly at least 10 % by weight in order to sufficiently improve the discoloration

In the present invention, if necessary, a wax such as a polyoleline wax or a paraffin wax can be addust to the resin binder. It is preferable that the amount of the wax is at least 0.1 part (part by weight, hereinatter referred to the same), particularly at least 0.5 part based upon 100 parts of the resm binder from the viewpoint of sufficiently imparting the effect of using the wax to the resin binder. However, when the amount of the wax is excessive, since there is a tendency that a film of the wax is formed on the electric lateril image of a photosensitive body, it is preferable that the amount of the wax is at most 20 parts, particularly at most 10 parts based upon 100 parts of the resin binder. Among the waxes, the polyolotine wax is 2 X

When a printed image is formed from the decolorizable toner containing the polycietins wax on a support such as copying paper, rough decolorization is scarcely caused and the punted unage can be rapidly and uniformly decolorized, even if the printed image is irradiated with near infrared rays from ority

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The reason why the decolorizable toner has such excellent properties is not evident. However, the following reasons can be thought as some parts of reasons.

When the decolorizable toner is fixed on a support, some particles of the wax are existed in the decolorizable toner, and the other particles of the wax are bled out between the particles of decolorizable toner and in the interface of the decolorizable toner and the support and on the outside surface of the decolorizable toner. The near infrared rays can be transmitted to the internal of the fixed decolorized toner diffusing effect. Furthermore, near infrared rays can also be transmitted to the upper surface, side surface and back surface of the near infrared ray - absorbing dye contained in the decolorizable toner owing to the owing to specific optical properties such as lens effect (effect such that the wax acts as a lens) and light light - reflecting function of the wax. Therefore, even if the decolorizable toner is irradiated with near intrared rays from only one direction, the near infrared rays are diffused in the toner and the near infrared ray absorbing dye can be rapidly decolorized. 33 \$

Also, the wax is softened by the irradiation of near intrared rays or healing as a supplementary means. mobility of the near infrared ray -absorbing dye and the decolorizing agent is heighlened, and the contact frequency of the near infrared ray - absorbing dye and the docolorizing agent is increased. That is, the wax acts as a lubricant, and the near infrared ray - absorbing dye is sufficiently decolorized. ţ

Accordingly, even if a material through which near intrand rays hardly pass is contained in the decolorizable toner, the near intrared ray - absorbing dye can be uniformly and quickly decolorized

As a resin binder praterably used in the decolorizable toner, resins having no compatibility with a Concrete examples of these resin binders are, for instance, polystyrene resins represented by polystyrene polyoletine wax are used in order to obtain a sufficient tens effect of the polyoletine wax and the take and the like, polyester resins represented by polyester, unsaturated polyester and the like, an epoxy resin, (meth)acrylic resins represented by polymethyl methacrylate and the like, a silicone resin. a fluoriccarbนม resin, polyamide resins, polyvinyl alcohol resins, polyurothane resins, and the like. The present invention is not limited to the exemplified ones. These resin binders are generally used alune or in admixture thereof 20

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In the present specification, the meaning of the sontence "the polyoletine wax has no compatibility with the resin binder" is intended to reter to that the polyoletine wax and the resin binder are existed as a

misture thereof when the polypletine wax and the resin binder are observed by means of a microscope or such that the border of the polyoletine wax and the resin binder can be clearly observed like a so-called island -in -sea structure in which the resin binder constitutes sea and the polyoletine wax constitutes the like and a border of the both can be clearly observed. As its example, there can be cited. for instance,

In order that the polyoletine wax shows a lens effect in an obtained decolorizable toner, it is necessary that the potyoletine wax is coloriess. The term "coloriess" means "transparent", "colored transparent" or "white". In the present invention, it is particularly preferable that the wax is transparent. When the polyoletine wax has compatibility with the resin binder, since the polyotetine wax does not show the lens effect in an obtained decolorizable toner, it is necessary that the polyplefine was has no compatibility with

Typical examples of the polyoletine wax are, for instance, unmodified polyoletine wax. modified polyolefine wax prepared by carring out a block or graft copolymerization of olefine components, and the

As the otefine components used in the unmodified polyotefine wax and modified polyotefine wax, any of a homopolymer made of a single otefine monomer and a copolymer made of an otefine monomer and the instance, ethylene, propylene, butene - 1, pentene - 1,3 - methyl - 1 - butene, 3 - methyl - 2 - pentene and the other oletine monomers can be cited. As the other monomers which can be copolymerized with the olefine monomer, for instance, various kinds of monomers such as the other olefine monomer, vinyt ethers other monomer copotymerizable with the oletine monomer can be used. As the oletine monomer, for such as vinyl methyl ether, vinyl esters such as vinyl acetate, haloolefines such as vinyl fluoride. (meth) acrylic acid esters such as methyl acrylate and methyl methacrylate, acrylic acid derivatives such as acrylonitrile, and organic acids such as acrylic acid can be cited. When the copolymer is used, copolymers such as an eithylene - propylene copolymer, an eithylene - butene copolymer, an eithylene - vinyl acetate copolymer, an ethylene-vinyt methyl ether copolymer and an ethylene-propylene-vinyt acetate copolymer can be cited. When the copolymer is obtained from the monomers other than the olefine monomer, it is preferred that the content of the otefine units derived from the otefine monomer is at least 50

As the component for modifying which is contained in the modilied polyoletine wax, for instance, butylstyrene; a -methylene aliphatic monocarboxytic acid ester monomers such as methyl acrylate, ethyl acrylate, methyl methacrylate and ethyl methacrylate, and the like can be used. When the aromatic vinyl particularly 1 to 10 % by weight. Also, when the a -methylene aliphatic monocarboxylic acid ester is used the component for modifying, it is preferred that its content is 0.1 to 50 % by weight, particularly 1 to 40 aromatic viny! monomers such as 1 - phenytpropene, styrene, methylstyrene, p - ethylstyrene and p - n monomer is used as the component for modifying, it is preferred that its content is 0.1 to 15 % by weight.

In the present invention, in addition to the above - mentioned ones, there can be used, for instance, a polyeithylene wax comprising a copolymer such as a block copolymer of ethylene and melhyl methacrylate or butyl methacrylate, a graft copolymer prepared by carrying out a graft polymerization of polyethylene and methyl methacrylate or butyl methacrylate, or a block copolymer of ethylene and styrene, and the like.

It is desired that the polyoletine wax itself has a low softening point. For instance, it is desired that the sottening point of the polyoletine wax measured by a ring and ball method prescribed in JIS K-2531 than the above-mentioned range, there is a tendency that particles of the wax are not remained as particles at the time an obtained decolorizable toner is fixed on a support, and therefore, lens effects cannot be sufficiently obtained. When the softening point is higher than the above - mentioned range, there is a therefore the wax hardly bleeds out between the particles of the decolorizable toner and in the interface of (1960) is 80° to 180° C, particularly 90° to 165° C. When the softening point of the polyolefine way is tower lendency that the wax is hardly molten when the obtained decolorizable toner is fixed on a support, and the decolorizable toner and the support and the like, and near infrared rays cannot be sufficiently

Also. It is preferred that the melt viscosity of the potypletine wax, which is measured by using a BL type

viscosimeter, is 20 to 6000 centipoise, particularly 50 to 4500 centipoise at 160 °C. When the mell viscosity is lower than the above - mentioned range, the wax is not existed in the state of particles at the time an obtained decolorizable toner is fixed on the support, and there is a tendency that tens effects cannot be sufficiently obtained. When the melt viscosity is higher than the above - mentioned range, the wax is hardly blect out between the particles of the decolorizable toner and in the interface of the decolorizable toner and the support and the like at the time an obtained decolorizable toner is leed on a support, and there is a tenclency that noar infrared rays cannot be sufficiently transmitted to the inside of the toner

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Concrete examples of the polyotefine wax are, for instance, VISCOL 660P and VISCOL 550P which are 320P. HI - WAX 220P. HI - WAX 2203A and HI - WAX 4202E which are commercially available from MITSUI commercially available from SANYO CHEMICAL INDUSTRIES. LTD. POLYETHYLENE 6A commercially available from ALLIED CHEMICAL CORP. HI-WAX 400P. HI-WAX 100P. HI-WAX 200P. HI-WAX WAX PE190 which are commercially available from HOECHST JAPAN LIMITED and the like, and the PETROCHEMICAL INDUSTRIES. LTD., HOECHST WAX PES20, HOECHST WAX PE130 and HOECHST present invention is not limited to the exemplified ones.

It is preferred that the average particle diameter of the polyotetine wax, which is measured by using a particle diameter NICOMP 270 imported by NOZAKI AND CO.. LTD. in accordance with a light scattering is smaller than the above - mentioned range, there is a tendency that the effect of bleeding out of the method, is 0.5 to 3 um, particularly 0.8 to 2 um. When the average particle diameter of the polyotetine wax polyolefine wax becomes insufficient. When the average particle diameter of the polyolefine wax is larger than the above -mentioned range, there is a tendency that a polycletine wax film is formed on an etectric latent image of the photosensitive body. 6 5

absorbing dye selected from the group consisting of a near infrared ray - absorbing dye represented by the In the present invention, as the near infrared ray -absorbing dye, at least one near infrared ray -

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wherein X⁻ is a halogen ion, perchloric acid ion, PF₆⁻, SbF₆⁻, OH⁻, sulfonic acid ion or BF₆⁻, Y⁻ is a cation having absorptions in the near intrared region, and a near intrared ray - absorbing dye represented

wherein each of R¹, R², R³ and R⁴ is independently hydrogen atom, a hydrocarbon group or a hydrocarbon group containing a hetero atom, Y* is the same as defined above, is used.

FCH;SO; - F2CHSO; - F3CSO; CICH;SO; - C3CHSO; - C3CSO; - CH;OCH;SO; - and (CH;)-3NCH;SO; - phenylsullonic acid ion such as C4H;SO; - substituted phenylsullonic acid ions such as As the hatogen ion, fluorine ion, chlorine ion bromine ion and iodine ion can be cited. As the sulfonic acid ion. methylsulfonic acid ion such as CH₃SO₂-; substituted methylsulfonic acid ions such as CHG.H.SO3- (CH3, RG.H.SO3- (CH3) RG.H.SO3- (HO) RG.H.SO3- (HO) RG.H.SO3- CH.CISO3- GH.CISO3- GH.CISO3- GH.CISO3- GH.CISO3- GH.CISO3- GH.F.SO3- GH. 8 \$

In the general formula (II), as the concrete examples of R¹, R², R³ and R⁴, there can be cited, for instance. hydrogen atom an alkyt group, an aryt group. an allyl group, an aralkyl group, an alkenyl group, an alkynył group, siłył group, a helerocyclic group, a substituted alkyl group, a substituted aryl group, a substituted allyt group, a substituted aralkyl group, a substituted alkenyl group, a substituted alkynyl group. a substituted stiyl group and the like. Among them, as the concrete examples of preferable ones, there can be cited. for instance, hydrogen atim, phenyl group, anisyl group, n - butyl group, n - pentyl group, n hexyl group. n-heptyl group. n-octyl group, n-dodecyl group. cyclohexyl group. cyclohexenyl group. methoxymethyl group, methoxyethyl group, ethoxyphenyl group, toluyl group, 1-bulylphenyl group, lluorophenyl group, chlorophenyl group, diethylaminophenyl group, vinyl group, allyl group, triphenylsityl graup, dimethylphenylsilyl graup, dibutylphenylsilyl graup, trimethylsilyl graup, piperidyl graup, thienyl group, furyl group and the like. It is preferred that at least one of R'. R? A? and R* is an alkyl group having butyl group. n – pentyl group. n – hevyl group. n – heptyl group. n – octyl group and n – dodecyl group are 1 to 12 carbon atoms. Among these alkyl groups, alkyl groups having 4 to 12 carbon atoms such as n -

and pyrylium. having absorptions in the near intrared regions, and the like can be preferably used. Typical As the preferable Y' of the near infrared ray - absorbing dye represented by the general formula (I). for instance, cationic dyes of cyaning. Liarytmethane, aminium, diimonium, thiazine, xanthene, oxazine, styryt examples of Y* are, for instance,

$$(C_2H_5)_2 N$$

$$C = CH - CH = CH - C$$

$$(C_2H_5)_2 N$$

$$(C_2H_5)_2 N$$

$$C_{2}H_{5}$$

$$C_{2}H_{5}$$

$$C_{2}H_{5}$$

$$C_{2}H_{3}$$

$$C_{1}H_{3}$$

$$C_{2}H_{3}$$

$$C_{2}H_{3}$$

$$C_{2}H_{3}$$

$$C_{2}H_{3}$$

$$C_{2}H_{3}$$

å

8

$$^{2}H_{5})_{2}N$$
 $^{2}C_{2}H_{5}$
 $^{2}C_{2}H_{5}$
 $^{2}C_{2}H_{5}$
 $^{3}C_{2}H_{5}$
 $^{3}C_{2}H_{5}$
 $^{3}C_{2}H_{5}$

and the like

The amount of the near infrared ray - absorbing dye is 0.01 to 25 parts, preferably 0.1 to 15 parts based upon 100 parts of the rein binder. When the amount of the near intrared ray - absorbing dye is less than the above - mentioned range, an obtained decolorizable toner cannot be sufficiently colored. When the amount of the near infraind ray - absorbing dye is more than the above - mentioned range, an amount of tubo - etectric charge of an obtained decolorizable toner is sometimes impaired.

In the present invention, as the decolorizing agent, a decolorizing agent represented by the general

ainm, an alkyt group, an aryt group, an altyt group, an aralkyt group, an alkenyt group, an alkynyt group, a hererocyclic group, a substituted alkyl group, a substituted aryl group, a substituted allyl group, a group, an alkenyl group, an alkynyl group, silyl group, a heterocyctic group, a substituted alkyl group, a is an alkyl group having 1 to 12 carbon atoms; and each of R? R10, R11 and R12 is independently hydrogen wherein each of R¹, R⁴, R¹ and Rª is independently an alkyl group, an aryl group, an allyl group, an aralkyl substituted alkynyl group or a substituted silyl group, with the proviso that at least one of R*, R*, R*, and R* substituted any group, a substituted allyt group, a substituted aralkyl group, a substituted alkenyl group. substitutert aralkyt group, a substituted alkenyt group or a substituted alkynyt group, is used.

dodecytriphenylborate, timethylhydrogenammonium n - butyttriphenylborate, triethylhydrogenammonium hvrate, tetramethylammonium n - octyltrianisythorate, tetraethylammonium n - bulyltriphenylborate. teirabutythorate, tetraethylaminonium tetrabutythorate, tetra - n - butylammonium tetra - n - butylborate, letramethylammonium tri – n – butylltriphenylsılyilborate. tetraethylammonium tri – n – butylltriphenylsilyl) – Concrete examples of the decolorizing agent are, for instance, tetramethylammonium n-butylretraethylammonium n.-butyltrianisylborate, tetrabutylammonium n.-butyltriphenylborate, tetrabutylam.nonium n - bulyltrianisylborate, tetraocilylammonium n - octytriphenylborate, tetrabutylammonium n tetramethylammonium tri - n - butyl tetrabutylam iriphenylborate, tetramethylammonium n - bulytirianisylborate, tetramethylammonium n - octyltriphenyl -(rirmathytphonylsilyl)borate. Tetraethylammonium tri - n - butyt(dimethylphenylsilyl)borate. tetrabutylammonin n - bulyl(triphenylsılyl)borate. tetramethylammonium n - butyttriphenylborate, tetrahydrogenammonium n - butylinphenylborate,

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dimethytphenyt(trimethylsityt)borate, and the like, and these decolorizing agents can be used alone or in tetrabutylammonium phenyisilyi)borate. tetraethylammonium n-octyldiphenyl(di-n-butylphenylsilyi)borate. tetrabulylamtetramethylammonium n - octyldiphenyl(di - n - butyl n - octyldiphenyt(di - n - bulylphenylsılyl)borate, tetramethylammonium dimethylphenyl(trimethylsilyl)borate. monium tri - n - butyt(dimethytphenylsilyt)borate. (trimethylsilyl)borate, tetraethylammonium monium

mentioned range, the speed of decolorization becomes slow. When the amount of the decolorizing agent is more than the above-mentioned range, light stability of a ligure or image formed from an obtained depends upon the kind of the near infrared ray - absorbing dye, but the amount of the decolorizing agent is usually adjusted within the range of 1 to 2500 parts, preferably 5 to 1000 parts based upon 100 parts of the near infrared ray - absorbing dye. When the amount of the decolorizing agent is less than the above -The amount of the decolorizing agent cannot be indiscriminately determined because decolorizable toner is deteriorated and the color of the figure or image is discolored or laded.

absorbing dye contained in the decolorizable toner by the irradiation of natural light, a tight fastness In the present invention, as mentioned above, in order to prevent discoloration of the near infrared ray -

As the light fastness stabilizer, for instance, a heat -resistant age resistor, a metal soap or a metal

As the heat -resistant age resistor, it is desired that a heat-resistant age resistor, which would not stain a support such as paper when the decolorizable toner is transferred to and fixed on the support, is oxide can be used. 8

Furthermore, when the decolorizable toner is applied to paper having a whity - color such as copying paper, a heat - resistant age resistor having a white or light color is preferably used.

alkytated phenol, bisphenol A, bisphenol S, ethyl p - hydroxybenzoate, n - propyl gallate, tauryl gallate and As the heat - resistant age resistor, a heat - resistant age resistor which scarcely stains a white copying hydroquinons and hydroquinone monosthyl ether; alkylated phenol and phenol derivative age resistors such paper and the like can be used. Concrete examples of the heat-resistant age resistor are, for instance. as 1-oxy-3-methyl-4-isopropylbenzene, 2.6-di-t-butylphenol, 2.6-di-t-butyl-4-ethylphenol. t - butyl - a - dimethylamino - p - cresol, 2 - († - methylcyclohexyt) - 4,6 - dimethylphenol, styrenated phenol hydroquinone derivative age resistors such as 2.5-di-t-amythydroquinone, 2.5-di-t-butyl-2.6 - di - 1 - dutyl - 4 - methylphenol. 2.6 - di - 1 - butyl - 4 - sec - butylphenol. butylhydraxyanisole, 2.6 - di -8 22

alone or in admixture thereof. Among these heat - resistant age resistors, the hydroquinone derivative and phenol derivative are preferable since they are excellent in compatibility with the styrenic resin binder and phenol). n -octadecy! – 3 - $(4^{\circ}$ - hydr(xy – 3.5° - di – 1 - butylphenyl)propionate: phosphite – age resistors such as tris(nonylphenyl) phosphile, tris(mixed mono and di-nonylphenyl) phosphile, phenyldiisodecyl phosphite, diphenylmono(2 - ethylhexyt) phosphite, diphenylmonotridecyt phosphite, diphenylisodecyl phosphite, diphenylisocctyl phosphite, diphenylphenyl phosphite, triphenyl phosphite, tristtridecyl) phosphite and letraphenyldipropylene glycol phrisphile, and the like. These heat - resistant age resistors can be used resorcinol; hindered phenol - age reistors such as 1.1.3 - tris - (2 - methyl - 4 - hydroxy - 5 - 1 - butyl phenyl)bulane, 4.4' – butylidene bis – (3 – methyl – θ – t – butylphenol), 2.2 – thiobis(4 – methyl – θ – t – butyl can remarkably prevent the deterioration of properties of a crystalline resin due to lights. 8

cannol be sufficiently prevented. When the amount of the heat-resistant age resistor is more than the resistant age resistor, and also the decolorizable toner cannot be sufficiently colored by the near infrared The amount of the heat - resistant age resistor is 0.05 to 30 parts, preferably 0.5 to 10 parts, particularly age resistor is less than the above - mentioned range, discoloration of the near infrared ray - absorbing dye preferably 0.5 to 2 parts based upon 100 parts of the resin binder. When the amount of the heal - resistant above - mentioned range, cost increases in accordance with the increase of the amount of the heat ţ

infrared ray - absorbing dye, as the light fastness stabilizer, a metal oxide or a metal soap can also be In the present invention in order to prevent the discoloration change of color or lading of the near contained in the decolorizable toner ŝ

Accordingly, it is thought that when the above - mentioned metal oxide or metal soap exists together with The reason why the metal oxide and metal soap impart discoloration resistance is not clear, but it is on its surface and the metal soap has an ionic polar group such as carboxyl group. That is, it is thought that since the near intrared ray – absorbing dye is an ionic complex, an ion pair of the complex is stable when an anionic polar group exists, and therefore, the stability of the dye to the light and heat is improved. thought that the discoloration resistance probably results from that the metal oxide has a basic polar group the near infrared ray = absorbing dyn, the near infrared ray = absorbing dye is stabilized, and the discolor = 55

ation, change of color and fading are prevented.

SiQ2+A1201, A1201, Na20+CO2, MgO+A1201, CO2 and the like. These metal oxides can be used alone or in Concrete examples of the metal oxide are, for instance. MgO. Al₂O₃, SiO₂, Na₂O. SiO₂ • MgO. admixture thereof. Among these metal oxides, MgO. a mixture of MgO and SiO2 or At2O3. Na2O. SiO2+MgO, SiO2+AI2O1, AI2O1, Na2O+CO2, MgO+AI2O1+CO2 and the like are preferable

When the metal oxide is used as a light fastness stabilizer, it is preferred that the amount of the metal oxide is 1 to 50 parts, particularly 5 to 20 parts based upon 100 parts of the resin binder. When the amount of the metal oxide is more that the above - mentioned range. There is a tendency that the color of an the above - mentioned range, there is a tendency that a sufficient effect for preventing discoloration cannot obtained toner is shaded by the color of the metal oxide. When the amount of the metal oxide is less than

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When the amount of the metal oxide is at least 5 parts based upon 100 parts of the resin binder, in formed portions are hardly distinguished from the other portions after decolorizing. Such advantage is Also, MgO, the metal oxide containing the MgO and a mixture thereof can be particularly preferably used in case an image is formed from the decolorizable toner of the present invention on a white copying paper which is usually used in electrophotography, and then the image is decolorized by the irradiation of near infrared rays, the decolorized portion of the image shows the same white color and gloss as the copying particularly noticeable when a metal oxide containing MgO is used among the light fastness stabilizers. paper because the gloss of resin binder is lowered. Therefore, there is an advantage that the image the present invention since the coloring of the dye is not hindered by them when an image is formed. 5

sometimes impaired, it is usually preferred that the average particle diameter of the metal oxide is at most 5 um, particularly at most 1 um. The shape and color of the particle are not particularly limited. However, in order to lower the gloss of the resin binder and traces which are formed by decolorizing the formed figure When the average particle diameter of the metal oxide is too large, since quality of the image is or image, it is preferable that the particle has a spherical or ellipsoidal shape, and that the color of the 2

Concrete examples of the metal soap are, for instance, salts of stearic acid such as lithium stearate, particle is white since the color of copying paper for electrophotography is generally white. ŗ

generated. When the amount of the metal soap is less than the above - mentioned range, there is a barium 2 - ethylhexylate, calcium stearate, magnesium stearate, calcium laurate, and the like are preferable because they have a melting point which is suitable for the toner and are not toxic. When the metal soap is used as the light fastness stabilizer, it is preferred that the amount of the metal soap is 0.05 to 10 parts. particularly 0.1 to 5 parts based upon 100 parts of the resin binder. When the amount of the metal scap is more than the above - mentioned range, there are tendencies that the amount of tribo - electric charge of the decolorizable toner is impaired, that wrong effects such that the metal soap adheres to a photosensilive body and the like are exhibited and that the photosensitive body are stained by the metal soap due to the bleeding out of the decolorizable toner to the surface, and as a result, defects of a formed image are anoate, zinc 2 - ethylhexanoate, cadmium 2 - ethythexanoate and tead 2 - ethylhexanoate; salts of ricinolic acid such as barium ricinoleate, zinc ricinoleate and cadmium ricinoleate; a dibasic lead salt of stearic acid such as 2PbO+Pb(C1, 1415 COO)2; salts of salicylic acid such as lead salicylate, zinc salicylate, tin salicylate and chromium salicylate; a tribasic lead salt of maleic add such as 3PbO+Pb(C4H2O4)H2O; a dibasic lead salt of phthalic acid such as 2PbO.Pb(CeH.O.), and the like. These metal soaps can be used alone or in admixture thereof. Among these metal soaps, zinc stearate, zinc laurate, lead salicylate, zinc ricinolate, magnosium stearate, aluminum stearate, calcium stearate, strontium, stearate, barium stearate, zinc stearate, cadmium stearate and lead stearate; salts of lauric acid such as cadmium laurate, zinc laurate, calcium laurate and barium laurate; salts of chlorostearic acid such as calcium chlorostearate. barium chlorostearate and cadmium chlorostearate; salts of 2-ethylhexanoic acid such as barium 2-ethylhexlendency that discoloration resistance and stability of images are not sufficiently improved. ŝ ç 8 38

ultraviolet adsorbing agunt which can effectively absorb ultraviolet rays having a wave length of 300 to 370 nm or so, which would easily yellow the resin binder, is preferably used. Concrete examples of the In the present invention in order to prevent the yellowing of the resin binder contained in the decolonizable toner, an ultraviolet adsorbing agent can be used. As the ultraviolet absorbing agent, an ultraviolet adsorbing agent are, for instance, benzophenone - ultraviolet adsorbing agents such as 2,4 dihydroxybenzophenone. 2 – hydroxy – 4 – methoxybenzophenone. 2 – hydroxy – 4 – octyloxybenzophenone. 2-hydroxy-4-dodecyloxybenzophenone, 2.2'-dihydroxy-4-methoxybenzophenone and 2.2'dihydroxy - 4,4" - dimethoxybenzophenone; salicyclic acid ester - ultraviolet adsorbing agents such as phe nyl salicylate, p - t - butylphenyl salicytate and p - octylphenyl salicylate; benzotriazole - ultraviolet adsorb ing agents such as 2 - (2' - hydroxy - 5' - methylphenyl)benzotriazole, 2 - (2' - hydroxy - 5' - t - bulyl phonyl)banzotnazote. 2 - (2' - hydroxy - 3'.5' - di - t - butylphenyl)benzotnazote, 2 - (2' - hydroxy - 3' - t -55

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2 - [2" - hydroxy - 3" - (3",4",5",6" - totrahydrophthalimidomethyt) - 5" methylphenyt|benzotriazole, a hindered amine - ultraviolet adsorbing ageirts such as bis(2.2.6.6 tetramethyl - 4 - piperidyl) sobacate, and the like. The present invention is not limited to libese exemplified 2 - (2' - hydroxy - 3',5' - di - 1 - turfylphenyl) - 5 ones. These ultraviolet adsorbing agents can be generally used aliane or in admixture thereof butyl - 5' - methylphenyl) - 5 - chlorobenzotnazole. and

It is preferred that the amount of the ultraviolet addorbing agent is 0.2 to 30 parts, particularly 0.5 to 5 adsorbing agent are not sufficiently obtained. When the amount of the ultraviolet adsorbing agent is more parts based upon 100 parts of the resin binder. When the amount of the ultraviolet adsorbing agent is less than the above - mentioned range, there is a tendency that effects exhibited by using the ultraviolat than the above -mentioned range, further improvements exhibited by using the ultraviolet adsorbing agenit are scarcely expected and there is a tendency that cost increases 5

If necessary, an adequate amount of additives such as a magnetic powder, a fluidizing agent, a plasticizer, a color pigment or a near infrared ray - reflecting or - absorbing material can be contained in the decolorizable toner of the present invention.

As the processes for preparing the decolorizable toner of the present invention, a solution process and a melting process can be employed. š

binder in an organic solvent mixing them, adding a decolorizing agent and a light fastiness stabilizer, and if necessary, adding an ultraviolet adsorbing agent, a wax or other additives thereto, dissolving and mixing them, removing the organic solvent from the obtained mixture, then, coasely pulverizing the mixture by using a hammer mill, cutter mill or the like and then finely pulverizing by using a jet mill or the like to give a The solution process is a process comprising dissolving a near infrared ray - absorbing dye and a resuldecolorizable toner having an average particle diameter of 5 to 30 um or so 2

absorbing dye and a resin binder, adding a decolorizing agent and a light fastness stabilizer, and if necessary, adding a wax, an ultraviolet absorbing agent, and other additives thereto, kneading linem, couling the obtained mixture, and then finely pulverizing the mixture in the same manner as the solution process to The melting process is a process comprising heating to melt and kneading a near infrared ray 2

resistance without the influence of heat history on the near infrared ray - absorbing dye during melting and ray - absorbing dye and a resin binder A having a softening point (a softening point according to a ring and ball method, hereinalter referred to the same) of at teast 30°C lower than the decomposition temperature of the near infrared ray - absorbing dye are heated between at least sottening point of the resin binder A and at least 10°C lower than the decomposition temperature of the near infrared ray - absorbing dye, molten and kneaded them, and to the obtained motten kneaded mixture is added a resin binder B having a softening point of at least 10°C higher than the softening point of the resin binder A and lower than the decomposition temperature of the near intrared ray - absorbing dye, and then the mixture is heated, inolleri In order to obtain a decolorizable toner which is not discolored or yellowed and has excellent offset kneading the near infrared ray - absorbing dye and the resin binder, it is preferred that the near infrared and kneaded for a short period of time. give a decolorizable toner. 8 8

In the process for preparing the decolorizable toner of the present invention, at first, the near infrared ray-absorbing dye and the resin binder A having a sottening point of at least 30°C lower than the decomposition temperature of the near infrared ray - absorbing dye are motten and kneaded. \$

polyvinyl chloride, polyvinyl acetate, polyethylene, polypropylene, polyester, polyurethane, polyariide, aii epoxy resin, polyvinyl bulyral, an acrylic acid resin, rosin, denatured rosin, a terpene resin, a phenol resin, a paraffin and a paraffin wax, which have softening points of at least 30°C lower than the octyl acrylate copolymer, a styrene-methyl methacrylate copolymer, a styrene-ethyl methacrylate copolymer, a styrene - butyl methacrylate copolymer, a styrene - methyl chloromethacrylate copolynner, a styrene - acrytonitrite copolymer, a styrene - vinyt methyl other copolymer, a styrene - vinyl ethyl ether isoprene copolymer, a styrene - acrylonitrile - indene copolymer, a styrene - maleic acid copolymer and a styrene - mateic acid ester copolymer, resins such as polymethyl methacrylate, polybulyl methacrylate, decomposition temperature of the near infrared ray - absorbing dye. These resin binders A can be generally Concrete examples of the resin binder A are, for instance, styrene resins such as polystyrene, poly p-chlorostyrene, polyvinylloluene, a styrene-p-chlorostyrene copolyrner, a styrene-propylone сорокутвг. а styrene – vinyltoluene сорокутег. а styrene - vinylnaphthalene copolymer. а styrene – methyl соројутви, а styrene - vinyl methyl ketone copolymer, а styrene - butadiene copolymer, а styrene acrylate copolymer, a styrene – ethyl acrylate copolymer, a styrene – butyl acrylate copolymer, a styrene – 25 ŝ ç

There is a necessity that the softening point of the resin binder A is at least 30°C lower than the decomposition temperature of the near infrared ray - absorbing dye. When the resin binder A does not have used alone or in admixture thereof.

shsorbing dye, since the softening point of the resin binder A is close to the decomposition lemperature of instory when the resin binder A and the near infrared ray - absorbing dye are molten and kneaded, and inereby properties of the near infrared ray - absorbing dye is deteriorated or the dye is decomposed. It is preferred that the softening point of the resin binder A is at least 40°C lower than the decomposition the near infrared ray - absorbing dye, the near infrared ray - absorbing dye is badly influenced by its heat the softening point of at least 30°C lower than the decomposition temperature of the near intrared ray emperature of the near infrared ray - absorbing dye.

The ratio of the resin binder A and the near infrared ray - absorbing dye is determined in accordance with the ratio of the resin binder A and the resin binder B and the amount of the near infrared ray absorbing dye against the total amount of the resin binder A and resin binder B.

binder A and near infrared ray - absorbing dye when the resin binder B is not used, but the resin binder A In the present invention, it is preferred that the decoloring agent is previously mixed with the resin and the near infrared ray - absorbing dye are motten and kneaded, from the viewpoint of increasing the contact frequency of the near infrared ray - absorbing dye and the decolorizing agent and sufficiently improving the decolorizing property during the irradiation of near infrared rays.

the amount of the decolorizing agent cannot be absolutely determined. However, as mentioned above, the amount of the decolorizing agent is generally adjusted within the range of 1 to 2500 parts, preferably 5 to Since the amount of the decoloring agent depends upon kinds of the near infrared ray - absorbing dye, 1000 parts based upon 100 parts of the near infrared ray - absorbing dye.

colorized and its properties deteriorate. It is preferable that the upper limit of the heating temperature is at most a temperature which is 70°C higher, particularly 30°C higher than the softening point of the resin When the resin binder A and the near infrared ray - absorbing dye are molten and kneaded, the heating iemperature is adjusted to at least the softening point of the resin binder A and at least 10°C tower than the than the softening point of the resin binder A, the resin binder A and the near infrared ray - absorbing dye non temperature of the near infrared ray - absorbing dye, the near infrared ray - absorbing dye is dis binder A from the viewpoint of the dispersibility of the near infrared ray-absorbing dye with the resin decomposition temperature of the near infrared ray - absorbing dye. When the heating temperature is lower cannot be uniformly mixed. When the heating temperature is not at least 10°C lower than the decomposi -

binder B are uniformly kneaded for a short period of time. Therefore, in the present invention, it is preferred After melting and kneading the near infrared ray - absorbing dye and the resin binder A, the resin binder B is added to the obtained molten kneaded mixture. When the resin binder B is added to the molten kneaded mixture which is in the state of melting, it is difficult that the molten kneaded mixture and the resin that the molten kneaded mixture is cooled and then pulverized, and the resin binder B is blended with the pulverized mixture, heated to melt and kneaded, or that the resin binder B is previously molten, and the molten kneaded mixture is added thereto and kneaded.

When the molten headed mixture is cooled and pulverized, it is preferred that the particle diameter of obtained pulverized mixture is adjusted to at most 5 mm or so from the viewpoint of uniformly dispersing the particles in the resin binder B.

It is preferred that the ratio of the resin binder A and the resin binder B, that is, the weight ratio of the than the above - mentioned range, dispersibility of the near infrared ray - absorbing dye with the resin binder B is lowered. When the ratio is exceeds the above-mentioned range, offset resistance of an resin binder Athe resin binder B is 10/90 to 90/10, particularly 30/70 to 70/30. When this ratio is smaller obtained decolorizable toner is lowered.

A and the resin binder B. When the amount of the near infrared ray - absorbing dye is smaller than the absorbing dye becomes insulficient. When the amount of the near infrared ray - absorbind dye exceeds the As mentioned above, it is preferred that the amount of the near infrared ray - absorbing dye is adjusted above - mentioned range, there is a lendency that dispersibility of the near infrared ray - absorbing dye with to 0.01 to 25 parts, particularly 0.1 to 15 parts based upon 100 parts of the total amount of the resin binder -mentioned range, there is a tendency that coloring property based upon the near infrared ray the resin binders A and B is lowered.

As the resin binder B, resins which can be used as the resin binder A are exemplified. Among these resins, a resin having a softening point of at least 10°C higher than the softening point of the resin binder A and lower than the decomposition temperature of the near infrared ray - absorbind dye can be used. When the resur binder B does not have a softening point of at least 10°C higher than the softening point of the of the resin binder B is at loast the decomposition temperature of the near infrared ray - absorbing dye, the man intrained ray - absorbing dye comes to be decomposed, and the decolorizable toner is not sufficiently resin binder A, there is a tendency that offset resistance becomes not insufficient. When the softening point

colored by the dye and is discolored.

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When the molten kneaded mixture and the resin binder B are molten and kneaded, the temperature of its mixture is adjusted within a range of at feast the softening point of the resui binder B and lower than the decomposition temperature of the near infrared ray - absorbing dye. It is preferred that the above mentioned temperature is at most a temperature of 50°C higher, particularly 30°C higher than the softening point of the resin binder B from the viewpoint of uniformly dispersing the molten kneaded mixture

After the temperature of the mixture of the molten kneaded mixture and the resin binder B is attained to the sottening point of the resin binder B, it is desired that a period of time for melting and kneading the preferably within 3 minutes, more preferably within 1 minute from the viewpoint of preventing the molten kneaded mixture and the resin binder B is as short as possible, and is generally in the resin binder B. 5

As mentioned above, since there is a necessity that the molten kneaded mixture is uniformly dispersed in the resin binder B for a short period of time, it is preferred that means for kneading such as a pressure decomposition and deterioration of the near infrared ray - absorbing dye. kneader, an extruder or a roll is used when they are kneaded.

classified by using a wind-force classifier or the like to obtain a decolorizable toner having a desired After kneading of the molten kneaded mixture and the resin binder B. the mixture is usually allowed to usually to room temperature After the cooled mixture is coasely pulverized by a hammer mill, a cutter mill or the like, the pulverized mixture is further pulverized by, for instance, a jet mill or the like, and then cool or cooled by using a suitable cooling means to a temperature at which a motten material is not existed. 20 5

The thus obtained decolorizable toner containing the near infrared ray - absorbing dye has excellent coloring property. excellent offset resistance, excellent discoloration resistance excellent blocking resistance average particle diameter of, for instance, 12 um or so.

a resin binder and a near infrared ray - absorbing dye and a component for decolorizing containing the decolorizing agent is employed as a process for producing the decolorizable toner of the present invention there can be dissolved the problems such that when the resin binder. The near infrared ray - absorbing dye and the decolorizing agent are heated and molten at a time the near intrared ray - absorbing dye is In case that a process for producing a decolorizable toner, comprising using a master batch containing contacted with the decolorizing agent, and thereby the near infrared ray - absorbing dye is decolorized and the like since the near infrared ray - absorbing dye is not influenced in its heat history. 8 23

batch itself is excellent in light stability and shows excellent discoloration resistance for a long period of mentioned above, since the decolorizing agent is not contained in the master batch, the master

The resin binder and the near infrared ray - absorbing dye and if necessary, a light lastness stabilizer are contained in the master bath. 33

In the master batch, the aforementioned resin binder, the aforementioned near infrared ray - absorbing dye and the aforementioned light fastness stabilizer can be used.

at most 200 parts based upon 100 parts of the resin binder used in the master batch. When the amount of the near infrared ray - absorbing dye is excessive, there is a tendency that it is difficult that the near It is preferred that the amount of the near infrared ray - absorbing dye is at most 300 parts, particularly \$

As the processes for preparing the master batch, a solution process and a melting process can be infrared ray - absorbing dye is uniformly dispersed in the resin binder.

removing the organic solvent or water under a reduced pressure to give a mass of the mixture, and The solution process comprises the steps of mixing a first solution obtained by dissolving a resin binder in an organic solvent with a second solution obtained by dissolving or dispersing a near intrared rayabsorbing dye, a light fastness stabilizer and it necessary a dispersant in the organic solvent or water pulverizing the mass by using a ball mill or the like to give a master batch. \$

aqueous solvents such as water and aqueous solution of ethanol, and the tike. The organic solvent used in ketones such as acetone and methyl ethyl ketone; ethers such as diethyl ether, veratrole and zene and toluene: halogenated hydrocarbon solvents such as dichloromethane and carbon tetrachloride. Concrete examples of the organic solvent are, for instance, alcohols such as ethanol and isopropanol: tetrahydrofulan; phenols such as phenol and cresol; other aromatic solvents such as benzene, bromoben the first solution can be the same as or different from the organic solvent used in the second solution. 20

In the first solution, it is preferable that the amount of the organic solvent is 50 to 800 parts, particularly 500 parts based upon 100 parts of the resin binder. When the amount of the organic solvent is less than the above - mentioned range, there is a tendency that the resin binder is not sometimes sufficiently 55

dissolved in the solvent. When the amount of the organic solvent is more than the above - mentioned range, there is a tendency that excessive cost is sometimes necessitated for removing the organic solvent and

solution there is no necessity that the near infrared ray - absorbing dye or the tight fastness stabilizer is In the second solution, it is preferable that the amount of the organic solvent is 10 to 5000 parts, particularly 20 to 3000 parts based upon 100 parts of the near infrared ray - absorbing dye. When the When the amount of the organic solvent is more than the above - mentioned range, there is a tendency that excessive cost is sometimes necessitated for removing the organic solvent and water, in the second dissolved in the organic solvent. If necessary, the near infrared ray - absorbing dye and the light fastness amount of the organic solvent is less than the above - mentioned range, there is a lendency that the near infrared ray – absorbing dye and the resin binder cannot be sometimes sufficiently dissolved in the solvent. stabilizer can be dispersed in the organic solvent or water in a pulverized state. 9

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infrared ray - absorbing dye, and a light fastness stabilizer by using a kneading machine such as a biaxial The above - mentioned melting process comprises heating to melt and kneading a resin binder, a near screw extruder and a kneader to obtain a master batch, or cooling the obtained kneaded mixture, then pulverizing the mixture in the same manner as in the above - mentioned solution process to give a master Š

The decolorizable toner contains the master batch and a decolorizing agent, if necessary, a resin binder, a wax and a component for decolorizing containing a white filler.

It is preferable that the amount of the resin binder used in the component for decolorizing is at least 50 parts, particularly at least 100 parts based upon 100 parts of the decolorizing agent. 2

llower, magnesium oxide, magnesium hydroxide, clay, line powder of sitica, and the like. These white fillers Concrete examples of the white filler are, for instance, titanium oxide, calcium carbonate, alumina, zinc can be used alone or in admixture thereol. Among these white fillers, titanium oxide, calcium carbonate, zinc flower and the like are preferable because these are excellent in coloring property 2

It is desired that the amount of the white filter is at least 0.5 part, preferably at least 2 parts based upon impart the effect exhibited by using the white filler. When the amount of the white filler is too much, there is a lendency that the color of the toner is laded. Therefore, it is desired that the amount of the white filter is al most 50 parts, preferably at most 30 parts based upon 100 parts of the resin binder for the master batch 100 parts of the rest binder for the master batch and the component for decolorizing in order to sufficiently and the component for decolorizing. 3

Also, in consideration of the dispersibility of the components of the master batch and the component for decolorizing is less than the above range, there is a tendency that the component for decolorizing is not uniformly dispersed in the master batch. When the amount of the component for decolorizing exceeds the above range, there is a tendency that the components of the master batch are not uniformly dispersed in decolorizing, it is desired that the amount of the component for decolorizing is 10 to 3000 parts, preferably 30 to 2500 parts based upon 100 parts of the master batch. When the amount of the component for the component for decolorizing. 3

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decolorizable toner in which the master batch is used is obtained by blending the master batch with the component for decolorizing, heating to melt and kneading them and cooling, and after that, coasely pulvelizing an obtained mass and further linely pulvelizing the mass with, for instance, a jet mill to give toner particles having an average particle diameter of 5 to 30 um or so and, if necessary, classifying. ş

irradiated with natural light during the storage of the master batch, the discoloration of the master batch can be provented and therefore, the master batch shows excellent storage stability. When a light fastness As mentioned above, since the decolorizing agent which promotes the discoloration of a near infrared discolored, and the master batch shows excellent thermal stability. Also, even though the master batch is stabilizer is contained in the master batch, the thermal stability and storage stability of the master batch are ray - absorbing dye is not contained in the master batch, the near infrared ray - absorbing dye is not ş

decolorizable toner having a sufficient amount of tribo-electric charge suitable for a developer of electropholography is obtained, an electric charge regulator or an electrically chargeable fine particle can be coated to the surface of a particle for the decolorizable toner. When the

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As a method for coating the electric charge regulator or the electrically chargeable fine particle on the surface of the particle for a decolorizable toner, dry - coating methods using a mechanicchemical reaction such as a high speed impact treatment method using a HYBRIDIZATION SYSTEM commercially available cially available from NIPPON PNEUMATIC MEG. CO. LTD. or the like, a mechanochemical surface fusing from NARA MACHINERY CO., LTD., a fluidized-healflow treatment method using, for instance, a NEW MALMELIZER commercially available from DALTON CORPORATION, a SURFUSING SYSTEM commer-

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MICRON CORPORATION or the like, and a treating method using a powder mixing machine such as a cially available from FREUND INDUSTRIES CO , LTD or the like, well-coaling methods such as a spray during method using a GRANULEX commercially available from FREUND INDUSTRIES CO., L.I.D. or the ENGINEERING CO., LTD. or the like, a method using a spray dryer, a method using a FLOW COATER the like are cited, but the present invention is not limited to the exemplified ones. Among these using, for instance, a MECHANOFUSING SYSTEM commercially available from HOSOKAWA super mixer; a spray coaling method using, for instance, a COATMIZER JETCOATING SYSTEM commun. like, a fluidized-bed drycoating method using a DISPACUAT communically available from NISSHIN commercially available from FREUND INDUSTRIES CO., LTD, and a method using a fluichzed - bud dryor, methods, the high speed impact treatment method is particularly preferable. and

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Usually, the amount of the electric charge regulator or electrically chargeable fine particle which is coated on the particle for a decolorizable toner is adjusted so that the amount of the tribo - electric charge (absolute value) of the particle for a decolorizable toner is 10 to 40 µC/g, particularly 15 to 30 µC/g. The amount of the electric charge regulator which is coated on the particle for a decolorizable toner is generally amount of the electrically chargeable fine particle is 1 to 50 parts, particularly 5 to 30 parts based upon 100 0 t to 10.0 parts, particularly 0 5 to 5 parts based upon 100 parts of the particle for decolorizable toner parts of the particle for a decolorizable toner 15

chargeable electric charge regulator can be used. These electric charge regulators can be generally used electric charge regulator, a positively chargeable electric charge regulator and a negatively alone or in admixture thereof. As the

Concrete examples of the positively chargeable electric charge regulator are, for instance, an electric charge regulator represented by the general formula (IV): 2

$$\begin{bmatrix} R^{1:s} & \\ R^{1:s} - R - R^{1:s} \\ \\ R^{1:s} \end{bmatrix} \stackrel{\bigoplus}{\bullet} (IV)$$

wherein R¹³ is an alkyl group having 1 to 8 carbon atoms or benzyl group, each of R¹⁴ and R¹⁵ is independently an alkyl group having 1 to 18 carbon atoms. Rth is an alkyl group having 1 to 18 carbon atoms or benzyl group, Z is hydroxyl group or amino group, n is 1 or 2; an electric charge regulator represented by the general formula (V):

$$\begin{bmatrix}
R_{14} - R - R^{40} \\
0 \\
0
\end{bmatrix}$$
(V)

quaternary aminonium salt resins, and the like. Typical examples of these positively chargeable electric charge regulators are for insaince, BONTRON P - 51 commercially available from ORIENT CHEMICAL INDUSTRIES, LTD, TP-415 commercially available from HODOGAYA CHEMICAL CO., LTD. FUJIKURA wherein each of R11, R11, R13 and R20 is independently hydrogen atom an alkyl group having 1 to 22 carbon atoms or an aralkyl group having 9 to 20 carbon atoms, A is a hoteropoly - acid ion, styrene - acrylic acid FCA - 201PB commercially available from FUJIKURA KASEI CO., L.TD., and the like ç 3

Concrete examples of the negatively chargeable elecrtic charge regulator are, for instance, an electric charge regulator represented by the general formula (VI): (VI)

wherein each of R²¹, R²², R²³ and R²⁴ is independently hydrogen atom, an alkyl group having 1 to 8 carbon atoms, an allyl group or a halogen atom, L is -S -, -SO₂ - or -C(R²³)(R²⁴) - wherein each of R²⁴ and R²⁴ is independently hydrogen atom or an alkyl group having 1 to 8 carbon atoms; styrene - sulfonic acid sall resins, and the like. Typical examples of these negatively chargeable electric charge regulators are, for 1001N commercially available from NIPPON KAYAKU CO., LTD., FUJIKURA FCA - The color of the second styrene is sufficient to the se

The color of the above - mentioned electric charge regulator is not particularly limited, but it is preferred that the electric charge regulator is transparent, white or faintly yellow so that the color of the decolorizable toner can be the same as the color of paper for electrophotography, which is usually white, and that traces of the decolorizable toner can be inconspicuous.

Concrete examples of the electrically chargeable fine particle are, for instance, fine particles of poly(meth)acrylic acid ester such as a fine particle of polymethyl methacrylate, a line particle of polymethyl
acrylate and a line particle of poly-2-ethylhexyl acrylate; electrically chargeable organic fine particles
such as a fine particle of polystyrene and a fine particle of polyvinylidene fluoride; and electrically
chargeable inorganic fine particles such as a fine particle of colloidal silica, a line particle of titanium
dioside, a line particle of molybdenum sulfide and a fine particle of high purity silica. These fine particles
are used alone or in admixture thereof

When the particle diameter of the above - mentioned electrically chargeable line particle is excessively larger than the particle diameter of the particle of the decolorizable loner, there is a lendency that the electrically chargeable line particle is hardly held on the surface of the particle of the decolorizable toner. Accordingly, it is generally preferred that the particle diameter of the electrically chargeable fine particle is at most about 10 %, particularly at most about 5 % of the particle diameter of the decolorizable toner.

After the thus obtained decolorizable toner of the present invention is printed and fixed on a support made of paper or the like, the printed portion can be decolorized by irradiating near infrared rays by using a semiconductor laser, a halogen tamp or a light emitting diode. After decolorizing the printed portion, printing can be repeatedly carried out on the decolorized portion.

Accordingly, the decolorizable loner of the present invention can be preferably used for printing on a support such as copying paper or a passenger ticket, which can be repealedly used by printing at the time of taking a train and erasing at the time of getting off, a coupon ticket and various tickets for admission.

The present invention is more specifically described and explained by means of the following Examples wherein all parts are by weight unless otherwise noted. It is to be understood that the present invention is not limited to the Examples, and various changes and modifications may be made in the invention without departing from the spirit and scope thereof.

Raw materials used in the following Examples, Reference Example and Preparation Examples are shown in Tables I to 6.

50	45	40	35	30	25	20	15	10	5
				EI.	Fable 1				
Raw material	Name				Contents	ts			
Resin binder	RE-1	Styrene-n-available	tyrene-n-butyl acrylate-n-butyl methacrylate e	TOATSU CHE	ethacrylate	copolymer (T)	B ALMATEX	Styrene-n-butyl acrylate-n-butyl methacrylate copolymer (TB ALMATEX CPR-200 commercially available from MITSUI TOATSU CHEMICALS INC.)	merclally
	RE-2	Styrene by TOATSU (tyrene butyl ocrylate cor TOATSU CHEMICALS INC.)	copolymer (1	IB ALMATEX	CPR-100 cor	nnici claffy a	Styrene-butyl ocrylate copolymer (TB ALMATEX CPR-100 commercially available from MILSUI TOATSU CHEMICALS INC.)	MIISU
	RE-3	Styrene-2- Carbide L	ityrene-2-ethylhexyl acrylate Carbide Industries Co., Inc.)	crytate copol , Inc.)	lymer (NIKA)	LITE NC-6100	commerciuli	Styrene-2-ethylhexyl acrylate colvolymer (NIKALITE NC-6100 commercially available from Niphon Cartide Industries Co., Inc.)	nongin mo
	RE-4	Polyester realn (r	restn (restn 1	naving light	yellow) (धि	-101 commerc	sially availab	Polyester resin (resin having light yellow) (IR-101 commercially available from Mitsubusin Rosen Co. 1141)	ibi edoi
	RE-6	Styrene In	from SANYO CHEMICAL INDUSTRIES, LTD.)	te-2-ethylhe INDUSTRIES	xyl acrylate s, LTD.)	copolymer (1	Ti-1850 com	Styren redutyl acrylate-2-ethylhaxyl acrylato copolymer ('l'i-1860 commercially available from Sanyo Chemical Industrius, LTD.)	iniste.
	RE-6	Styrene-36	rylonitiile a	spolymer (AS	-FD commen	cially available	e from SIO	Styrene-acrylonitrille copolymer (AS-FD commercially available from SiOWA DENKO K.K.)	G
	RE-7 RB-6	Polycater Polymethy	Polycater resin (NB-382 commercially available from Nao Corporation) Polymethyl methacrylate resin (SUMIPEX B 1.6-6 commercially available	2 commercial c resin (SUR	ily available APEX B 1.C-	from Kao Col	poration)	Polycater resin (NB-382 commercially available from Nao Corporation) Polymethyl methacrylate resin (SUMIPEX B 1.G-6 commercially available from Sumitomo Chemical	o Chemical
	9-86	Company,	Company, Limited)	orviate-n-bu	tyl acrylate	copolymer (a	selt vincosit	Company, Limited)	oise (110°C).
	1	3 × 10*	3 x 10° poise (140°C))					:	
	RE-10	Polyester Styrene-e	Polycater (melt viscosity: 9 x 10 poise (110°C), 8 x 10° poise (140°C)) Styreng-ethyl acrylate copolymer (melt viscosity: 9.5 x 10° poise (110°C)	ty: 9 x 10 copolymer (poise (110° melt viscosi	c), 8 x 10 ³ ty: 9.5 x 10 ⁶	poise (140°C) poise (110°)) C), 3 x 10"	Polyester (melt viscosity: 9×10^4 poise (110°C), 8×10^3 poise (140°C)) Styrene-ethyl acrylate copolymer (melt viscosity: 9.6×10^9 poise (110°C), 3×10^4 poise (140°C))
	RE-12	Styrene	-butyl scryls	te copolymer	r (inelt visc	Sity: 4 x 10	• poisc (110	C), 2 x 10°	Styrene-n-butyl scrylate copolymer (melt viscosity: 4 x 10° poise (110°C), 2 x 10° poise (140°C))
	RE-13	Styrene-a	nethyl ocrylat	te copolymes	(melt visco : (melt visc	sity: 5 x 10° 5slty: 7 x 10	Poise (110	C). 4 x 10*	Styrene-methyl ocrylate copolymer (melt viscosity: 5 x 10° poise (110°C), 4 x 10° poise (140°C)). Styrene-m-butyl ocrylate copolymer (melt viscosity: 7 x 10° poise (110°C), 4 x 10° poise (140°C))

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Raw material	Name	Contents
Resin binder	RB-15	Styrene n-butyl acrylate copolymer (light transmittance: 91 %)
	RE-16	Styrene-n-butyl methacrylate copolymer (light transmittance 83 %)
	RE-17	Polyester (light transmittance: 65 %)
	RE-18	Styrene-methyl methacrylate-n-butyl ocrylate terpolymer (light transmittance: 91 %)
	RE-19	Styrene-2-ethylhexyl acrylate-n-butyl acrylate terpolymer (light transmittance: 83 %)
	RE-20	Epoxy resin (light transmittance: 68 %)
	RE-21	Polyester (light transmittance 80 %)
	RE-22	Polyester (light transmittance 60 %)
	RE-23	Booxy resin (light transmittance 60 %)
	RE-24	Styrene-n-butyl scrulate coodynau (sector)
	RE-25	Styrene-1-ethylacrylate copolymer (content in weight ratio: 94:6, softening point: 100°C)
	RE-26	Styrene-2-ethylhexyl acrylate copolymer (content in weight ratio: 92:8, softening point: 106°C) Low molecular weight polyethylene (SANWAX 161-P commercially available from SANYO CHEMICAL. INDUSTRIES, LTD.)
1	Resin bino	
	REA-1	Styrone-n-butyl acrylate-n-butyl methacrylate copolymer (softening point: 100°C, glass transition temperature: 57°C)
	REA-2	Styrene-n-butyl acrylate-methyl methacrylate copolymer (softening point: 68°C, glass transition temperature: 59°C)
R	REA-3	Styrens n-butyl acrylate copolymer (softening point: 93°C, glass transition temperature: 60°C)
	lesin bind	er B
	REB-I	Styrene-n-butyl acrylate copolymer (softening point: 128°C, glass transition temperature: 65°C)
	REB-2	Styrene-2-ethylhexyl acrylate copolymer (softening point: 112°C, glass transition temperature: 65°C) temperature: 60°C)
	REB-3	Styrene-methyl methacrylate copolymer (softening point: 125°C, glass transition temperature: 65°C)

Table 2

Raw material	Name		Contents	
Near infrared ray-absorbing dye	DY-1	n-C ₄ U ₀ -B⊖-(⟨◯) ₃ .	$(C_2 II_5)_2 N$ $C = CII - CII = CII - C$ $(C_2 II_5)_2 N$ $(C_2 II_5)_2 N$	
	DY-2	(n-C ₄ lig) ₂ -B≘(⟨⟨⟩) ₂ .	$\begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \\ \\ \\ \end{array} \\ \end{array} \\ \begin{array}{c} \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \end{array} \\ \end{array} \\ \begin{array}{c} \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \end{array} \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \end{array} \\ \\ \begin{array}{c} \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \\ \\ \end{array} \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ $	
	DY-3	(u-C ⁶ [1 ¹³) ³ -B⊖-	CH3 CH-CH-CH3 CH3 CH3 CH3 CH3	
	DY-4	$n-C_{12}H_{26}-B^{\Theta}-(\bigcirc)_3$.	$(C_2 I_5)_2 N$ $C = CII - CII = CH - C$ $(C_2 I_5)_2 N$ $(C_2 I_5)_2 N$	

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Raw material	Name		Contents	
Near infrared ray-absorbing dye	DY-5	t ^e .	$C_{2} = C_{11} - (C_{11} = C_{11}) \xrightarrow{3} C_{2} = C_{2$	
·	DY-6	CII3 C6 II4 SO3 € .	$(C_2 II_5)_2 N$ $C=CII-CII=CII-C$ $R(C_2 II_5)_2$ $R(C_2 II_5)_2$	
	DY-7	B₽4 [⊕]	$(C_2 II_5)_2 N$ $C = CII - CII = CII - C$ $C = CII - CII = CII - C$	
	DY-8	n-C4∥9-B⊕(((())3 ·	$CI \xrightarrow{S} CII-CII \xrightarrow{C} CII-CII \xrightarrow{S} CI$	

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Continue			
Raw material	Name		Contents
Near infrared ray-absorbing dye	DY-9	n-c4119 -18 ⁰ -() 3	$Cil-(Cil = Cil)_{\frac{1}{3}} \bigcirc \frac{1}{c_2 l l_5}$
	DY-10	cn3c2n4203	$(C_2 II_5)_2 N$ $C = CII - CII = CII - C$ $N(C_2 II_5)_2$ $N(C_2 II_5)_2$

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Table 3

Raw material	Нато	Contents
Decolorizing	SE-1	Tetrabutylammonium n-butyltriphenyl bornte
agent	SE-2	Tetrabutylanmonium dimethyldlphenyl boruto
	SE-3	Tetrabutylammonium dibutyldiphenyl borate
	SE-4	Tetrametylammonium n-butyltriphenyl borate
	SE-5	Tetrabutylammonlum tetra-n-butyl borate
	SE-6	Triethylhydrogenammonium n-butyltriphenyi borate
Vax	WA-1	Polypropylene wax (VISCOL 550P commercially available from SANYO CHEMICAL INDUSTRIES, LTD.)
Polyolefine wax)	₩A-2	Polypropylene wax (VISCOL 330P commercially available from SANYO CHEMICAL INDUSTRIES, LTD., softening point: 152°C, melt viscosity: 4000 centipoise (160°C))
	WA-3	Polyethylene wax (III-WAX 720P commercially available from MITSUI PETROCHEMICAL INDUSTRIES,
		LTD. softening point: 118°C, melt viscosity: 3800 centipoise (160°C))
	WA-4	Polyethylene wax (III-WAX 110P commercially available from MITSUI PETROCHEMICAL INDUSTRIES.
		LTD, softening point: 113°C, melt viscosity: about 12 centinoise (160°C))

15 10 25 20 75

Table 4

Raw material	Name	Contents
Near infrared	ÀD−1	Heavy calcium carbonate (LIGHTON A commercially available from SHIRAISHI CALCIUM KAISYA, LTD.)
ray-reflecting	AD-2	Silica (SS-33S commercially available from Tokuyama Soda Co., Ltd.)
gnid-roeds—ro	AD-3	Titanium oxido (ECT-52 commercially available from TITAN KOGYO KABUSHIKI KAISYA)
material	AD-4	Certum oxide (MIREK commercially available from MITSUI MINING & SMELTING CO., LTD.)
	AD-6	Titanium oxide (CR-60 commercially available from ISIIIIIARA SANCYO KAISIIA, LTD.)
Light	Motal ox	ida
fastness	OA-1	MgO (purity: 98.4 %, average particle diameter: 0.06 µm)
štabilizer	OA-2	Al_BO_B (purity: 64.0 % average particle diameter: 0.5 μ m)
	QA-3	Mixture of MgO and Λl _m O _m (purity: 62.7 %, MgO/Λl _m O _m (weight ratio: 1/1, average particle diameter: 0.3 μ m)
	OA-4	Al ₂ O ₅ ·Na ₂ O·CO ₂ (Al ₂ O ₅ : 35 % by weight, Na ₂ O: 17.7 % by weight, CO ₂ : 24.4 % by weight, others such as CaO 22.9 % by weight, average particle diameter: 0.5 μ m)
	OA-6	MgO·Al _a O _a ·CO _a (MgO·38.2 % by weight, Al ₂ O _a : 16.1 % by weight, CO _a : 7.6 % by weight, others: 38.1 % by weight, overage particle diameter: 0.3 μ m)
	QA-6	SiO ₂ ·MgO (SiO ₂ : 64.9 % by weight, MgO: 13.6 % by weight, others: 21.6 % by weight, average particle diameter: 0.05 μ m)
	OA-7	SiO ₂ ·Al ₂ O ₃ (SiO ₃ : 60.2 % by weight, Λ l ₂ O ₃ : 10.5 % by weight, others: 24.6 % by weight, average particle diameter: 0.5 μ m)
	Metal so	ip
	MS-1	Zinc stearate (SZ#2000 commercially available from Sakoi Chemical Industry Co., Ltd.)
	MS-2	Zinc laurate (2-12 commercially available from Sakal Chemical Inchestry Co., Ltd.)
	MS-3	Barlum stearate (commercially available from WAKO JUNYAKU KOGYO KABUSHIKI KAISHA)
	MS-4	Calcium stearate (commercially available from WAKO JUNYAKU KOGYO KABUSIIIKI KAISHA)
	MS-5	Lead 2-ethylhexylate (commercially available from WAKO JUNYAKU KOGYO KABUSHIKI KAISHA)

Table 5

Raw material	Mame 	Contents
Light	llest-re	sistant age resistor
featness	۸٥-1	1-oxy-3-methyl-4-isopropylbenzene
stabilizer	AO-2	2,6-Dihydroquinone
	AO-3	Alkylated obenet (Apalaula a ann
	A0-4	Alkylated phenoi (Antioxidant NV3 commercially available from BASE AKITIENGESELLSCHAFT)
	AO-5	p-Diphenylamine
	AO-6	2,2-Bis(4-hydroxyphenyl)propane
	AO-7	Bthyl p-hydroxybenzoate (PEP-FP commercially available from UENO FINE CHEMICALS INDUSTRY, LTD.)
	AO-8	n-Propyl gallate
	AO-9	Lauryl gallate
	AO-10	Resorcing
	AO-11	3.4-Dihydroxy-4'-methyl diphenylsulphone (CD-180 commercially available from SIKWA DENKO K.K.)
erge	CR-1	Quaternary ammonium sait (BONTRON P-51 commercially available from ORIENT CHEMICAL INDUSTRIES
rulator	CR-2	KAYACHARCR N-1 COMMUNICAL INDUSTRIES
	CR-3	KAYACHARGS N-1 commercially available from NIPPON KAYAKU CO., LTD.
		FCA-1001N commercially available from FUJIKURA KASRI CO., LTD. TP-415 commercially available from IKODOGAYA CHEMICAL CO., LTD. FCA-201PB commercially available from IKODOGAYA CHEMICAL CO., LTD.
		TOM INDOGRAM CIRMICAL CO.
		FCA-201PB comercially available from FUJIKURA KASRI CO., LTD.
×	CR-5	Coally available from FUJIKURA KASRI CO., LTD.
۰.		FCA-201PB comercially available from FUJIKURA KASRI CO., LTD.
	CR-5	The Colly available from FUJIKURA KASRI CO., LTD.

Raw material	Name	Contents
Ultraviolet	UA-1	Phenyl salicylate
gnidroeds	UA-2	2-discharge
agent	UA-3	2-flydroxy-1-methoxybenzophenone
	UA-4	2-(2-llydroxy-6-methylphenyl)benzotrinzole Bis (2,2,6,6-tetramethyl-4-piperidyl)sebacate
Dectrically	Organic	fine particles
chargeable fine	TE-I	Polymethyl methacrylate (FINE POWDER MP1000 commercially available from SOKEN CHEMICAL &
	TB-2	Polymethyl methacrylate (RIMS powDER Amona). In negatively chargeable)
,	TE-3	Polyvinylidene fluoride (KYNAP 3018 annual 11 m. negatively chargeable)
	Inorganic	COMPANY, LTD., average particle diameter: 0.3 μ m, negatively chargeable)
	TB-4	Colloids silica (ARROSIL R-972 commercially available from NIPPON ARROSIL CO., LTD., average particle diameter: 0.1 μ m, negatively chargeable)
	TB-5	Titanium dioxide (IDEMITSI) TITANIA IT-00
	T2-6	I.To., average particle diameter: 0.2 μ m, negatively chargesize) Titanium dioxide (commercially available from TITAN KOCYO KABUSIKI KAISIA, average particle diameter: 0.2 μ m, negatively chargeable)
ganic	SO-1	Tetrahydrofuran
vent	SO-2	Toluene
	SO-3	Dichloromethane
	S0~4 ·	Ethyl acetato
		Methyl sthyl ketone

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55 Example 1

(Experiments 1 to 10 and Comparative Experiments 1 to 4)

Raw materials shown in Tables I to 3 and 5 were blended in a blending ratio shown in Table 7 to give a uniformly blended mixture.

Then, 20 parts of the mixture and 80 parts of 1,1,2,2 - tetrachloroethane were mixed together, and the mixture was sufficiently dissolved to or dispersed in the 1,1,2,2 - tetrachloroethane to give a toner solution.

The toner solution was coated on a white copying paper with a brush so that a toner layer having a thickness of 20 to 30 µm in dry could be formed, and a solvent was vaporized to remove from the toner layer to give a sample.

As a physical property of the sample, light resistance A was evaluated in accordance with the following method. The result is shown in Table 7.

(Light resistance A)

The sample was allowed to stand for 10 days at room temperature in the shade of a room which gets direct rays of the sun through a window glass during fine weather. The discoloration of the sample was examined by comparing the sample with a sample to which the natural light was not irradiated, and evaluated in accordance with the following criteria for evaluation.

[Criteria for evaluation]

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- A: No change of color is observed in the sample
 - B: A slight change of color is observed in the sample.
 - C: A little change of color is observed in the sample.
 - D: Change of color is obviously observed.

When the decolorizing property of the decolorizable toners obtained in Experiments 1 to 10 of Example

1 was examined by irradiating near infrared rays to the decolorizable toners, it was observed that all of the
decolorizable toners stood comparison with the toners not containing a heat resistant age resistor and
exhibited practically satisfactory decolorizing property.

50	45	95	3 0	eo 15	5	5
			Table 7	N		
Experiment		Components	Components of decolorizable toner (parts)	oner (parts)		
ġ Ż	Resin bluder	Near infrared ray-absorbing dye	Decolorizing agent	Wex	Light fastness stabilizer (Heat-resistant age resistar)	- Light resistance A of decolorizable toner
1		DY-1 (2)	SE-1 (2.84)	WA-1 (5)	AO-2 (2)	
N			SR-1 (2.84)	WA-1 (5)	AO-6 (20)	< «
n		DY-1 (2)			A0-3 (2)	: •
₹	RE-1 (100)		SE-1 (2.84)		A0-4 (6)	٠,
ø	RE-2 (100)	DY-3 (20)			(2) 1 20	٠ ،
9	RB-3 (100)	DY~4 (10)			40-4 (15)	< ۰
2		DY-5 (16)				< ⋅
0 0	RB-2 (100)					<
o,	RE-1 (100)	DY-7 (10)				< -
10	RE-2 (100)	DY-8 (2)		WA-1 (5)	(6) VO+ (6)	< <
Comparative						
A Des linear						
-	RE-1 (100)	DY-1 (2)	SE-1 (2.84)	WA-1 (5)	l	(
8	RB-2 (100)	DY-2 (2)	SE-2 (5)	(c) (c)		ပ
m	RB-3 (100)	DY-3 (2)	SP-3 (5)		ı	۵
•	RE-1 (100)	DY-4 (10)	SB-2 (10)	WA-1 (0,05)	1 1	U d

From the results shown in Table 7, it can be understood that all of the decolorizable toners containing a 55 heat resistant age resistor obtained in Experiments 1 to 10 have excellent light resistance.

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[Experiments 11 to 19 and Comparative Experiments 5 to 7]

The procedure of Experiments 1 to 10 and Comparative Experiments 1 to 4 of Example 1 was repeated to give a toner solution except that raw materials shown in Tables 1 to 3 and 5 were blended in a blending ratio shown in Table 8.

Then, the procedure of Experiments 1 to 10 and Comparative Experiments 1 to 4 of Example 1 was repealed to give a sample.

As physical properties of the sample, light resistance B and decolorizing property were evaluated in accordance with the following methods. The results are shown in Table 8.

(Light resistance B)

The sample was allowed to stand for one week under a fluoroescent lamp (illumination intensity: 1500 lux). A reflection density of the sample after allowing to stand and the reflection density of the sample before allowing to stand were measured by using a Macbeth densitometer, and the rate of change of reflection density was calculated by the following equation.

[Rate of change of reflection density (%)]

= ((Reflection density of the sample after allowing to stand)/(Reflection density of the sample before allowing to stand) x 100]

Then, the light resistance B was evaluated in accordance with the following criteria for evaluation.

[Criteria for evaluation]

- A: Rate of change of reflection density is not less than 80 % (Light resistance is very excellent).
- 8 Rate of change of reflection density is not less than 80 % and less than 80 % (Light resistance is excellent).
- C: Rate of change of reflection density is not less than 40 % and less than 60 % (Light resistance is good).
- D: Rate of change of reflection density is less than 40 % (Light resistance is bad).

When the decolorizing property of the decolorizable toners obtained in Experiments 11 to 19 of Example 1 was examined by irradiating near infrared rays to the decolorizable toners, it was observed that all of the decolorizable toners stood comparison with the toners not containing a heat resistant age resistor of Comparative Experiments 5 to 7 and exhibited practically satisfactory decolorizing property.

			Tal	Table 8		
Ryneriment		Components	Components of decolorizable toner (parts)	oner (parts)		
	Resin binder	Near infrared ray-absorbing dye	Decolorizing	Wax	Light fastness stabilizer (Heat-resistance age resistor)	Light resistance is of decolorizable
	RE-1 (100)	DY-1 (2)	SB-1 (3)	WA-1 (3)	VO-6 (I)	<u> </u>
	RE-1 (100)	DY-2 (2)	SG-1 (4)	WA-1 (3)	AO-6 (5)	п
	RE-3 (100)	DY-3 (2)	SE-2 (6)	WA-1 (3)	AO-7 (2)	B
	RG-3 (100)	54-4 (2)	SB-2 (3)	7A-1 (3)	AO-7 (6)	«
	RE-2 (100)	DY-6 (2)	SE-3 (3)	WA-1 (3)	AO-8 (0.6)	8
	RB-2 (100)	DY-6 (2)	SE-3 (4)	WA-1 (3)	AO-8 (2)	<
	RE-2 (100)	DY-1 (2)	SE-3 (4)	WA-1 (3)	(I) 6-0V	c
	RE-1 (100)	DY-1 (2)	SE-4 (4)	WA-1 (3)	VO-9 (5)	<
	RR-3 (100)	DY-1 (2)	SE-4 (3)	WA-1 (3)	AO-10 (2)	<
Comparative Experiment						
	RE-1 (100)	DY-1 (2)	SE-1 (3)	WA-1 (3)	1	Q
	RE-2 (100)	DY-2 (2)	SB-1 (4)	WA-1 (3)	1	٥
	RE-3 (100)	DY-3 (2)	SE-2 (5)	WA-1 (3)	ı	a

From the results shown in Table 8, it can be understood that all of the decolorizable toners containing a 55 heat - resistant age resistor obtained in Experiments 11 to 19 have excellent light resistance.

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Examp	θ	2
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[Experiments I to II and Comparative Experiments I to 2]

Raw materials shown in Tables 1 to 3 and 5 were blended in a blending ratio shown in Table 9 and weighed so that the total amount could be 800 g, and all of the raw materials were thrown into a pressure kneader having an effective volume of 2 t. Then, the kneader was heated, and the kneading temperature of the raw materials was adjusted to 130°C. After the raw materials were kneaded for 15 minutes, the obtained kneaded material was taken out from the kneader and cooled to give a solidified product.

Then, the obtained solidified product was ground with a jet mill and classified by using an air classifier to give toner particles having an average particle diameter of about 12 µm. To the toner particles was added 0.3 % by weight of silica commercially available from NIPPON AEROSIL CO., LTD. under the trade name of R = 972, and they were mixed by using a Henschel mixer to give a toner.

As physical properties of the toner, decolorizing property and visible light resistance were evaluated in accordance with the following methods. The results are shown in Table 9.

(Decolorizing property)

Four parts of the toner and 100 parts of a silicone resin - coated carrier commercially available from POWDERTECH CO., LTD. under the trade name of F97 - 2535 were mixed to give a uniformly mixed developer.

A copying machine commercially available from SANYO ELECTRIC CO., LTD. under the trade name of SFT-Z70 was charged with the obtained developer, and an image was duplicated on a paper for plain paper copy using a lest chart No. 1-R1975 prescribed by THE SOCIETY OF ELECTROGRAPHY OF 35 JAPAN.

Near intrared rays were irradiated to the image - duplicated paper from an aluminum - coated halogen tamp (2 W/cm²) which was placed at a distance of 30 cm from the paper for 3 seconds. Five sheets of the paper and five sheets of non - duplicated paper were put on a perpendicular wall at random, and they were observed with naked eyes whether they were duplicated or not at a distance of 5 m from the wall. The distinction was carried out by live men and live women, and when the duplicated paper was recognized as a non - duplicated paper, one point was given. Then, total point of ten persons was counted, and decolorizing property was evaluated in accordance with the following criteria for evaluation.

U.

(Criteria for evaluation)

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- A: 40 to 50 points
- B: 30 to 39 points
- C: 20 to 29 points
- D: 0 to 19 points

(Visible light resistance)

After a transparent glass vessel was charged with the obtained toner and the toner was preserved for 10 days at room temperature in an experimental room which could get natural light, the color of the toner was compared with the color of a toner which was preserved for 10 days at room temperature in a dark room with naked eyes, and visible light resistance was evaluated in accordance with the following criteria for evaluation.

[Criteria for evaluation]

- A: No change of color is observed in the toner.
- B: A slight change of color is observed in the toner, but there is practically no problem in the toner.
- C: A little change of color is observed in the toner, but there is practically no problem in the toner.
- Change of color is obviously observed in the toner, and the toner is not suited for practical uses.

				Table 9	(S)			
Experiment			Components of	Components of decolorizable toner (parts)	r (parts)		Physical properties of decolorizable toner	ties of ner
d	Resin binder	binder	Near infrared ray-absorbing dye	Decolorizing	Wax	Electric charge regulator	Decolorizing property	Visible light
	6-3Y	(100)	DY-1 (5)	SB-1 (3)	WA-1 (3)	CR-1 (3)	<	<
N	RE-10	(100)	DY-1 (2)	SE-1 (1)	WA-1 (3)	g-1 (3)	<	<
m	RB-11	(100)	DY-1 (3)	SE-1 (2)	WA-1 (3)	CR-1 (3)	=	<
•	RB-12		DY-1 (6)	SE-1 (4)	WA-1 (3)	OR-1 (3)	<	, E3
ø	RE-0	(300)	DY-2 (4)	SE-1 (3)	WA-1 (3)	OR-1 (3)	<	<
9	RE -9	(100)	DY-3 (6)	SE-5 (3)	WA-1 (3)	3	<	<
7	RB-9	(100)	DY-4 (3)	SE-3 (2)	WA-1 (3)	G-1 (3)	<	<
&	RE-9	(100)	DY-5 (5)	SR-6 (10)	WA-1 (3)	CR-1 (3)	<	<
G 1	GE -9	(100)	DY-6 (2)	SE-1 (4)	WA-1 (3)	CR-1 (3)	<	<
10	RE-9	(100)	DY-7 (4)	SR-5 (8)	WA-1 (3)		<	<
1.1	RE-9	(100)	DY-8 (6)	SR-3 (3)	WA-1 (3)	CR-1 (3)	<	<
Comparative							•	
	RE-13 (100	(100)	DY-1 (5)	SE-1 (3)	WA-1 (3)	CR-1 (3)	·	<
2	RE-14 (100	(001)	0.7-1 (5)	SP-1 (3)) 4	: 1

From the results shown in Table 9, it can be understood that all of the decolorizable toners containing a resin binder having a melt viscosity of not more than 10⁷ poise at 110°C and a melt viscosity of not less than 10³ poise at 140°C obtained in Experiments 1 to 11 have excellent decolorizing properly and excellent visible light resistance.

Example 3

[Experiments 1 to 10 and Comparative Experiments 1 to 5]

Raw materials shown in Tables 1 to 3 were weighed in a ratio shown in Table 10 so that the total amount could he 800 g, and all of the raw materials were thrown into a pressure kneader having an effective volume of 2 t. Then, the kneader was heated, and the kneading temperature of the raw materials was adjusted to 120 °C. After the raw materials were kneaded for 5 minutes, the obtained kneaded material was taken out from the kneader and cooled to give a solidified product.

Then, the obtained solidified product was ground with a jet mill and classified by using an air classifier to give toner particles having an average particle diameter of about 11 µm. To the toner particles was added 0.2 % by weight of hydrophobic silica commercially available from NIPPON AEROSIL CO., LTD, under the trade name of R - 972, and they were mixed by using a Henschel mixer to give a toner.

As a physical property of the toner, printed trace after decolorizing was evaluated in accordance with the following method. The results are shown in Table 10.

(Printed trace after decolorizing)

Four parts of the toner and 100 parts of a silicone resin - coated carrier commercially available from POWDERTECH CO., LTD, under the trade name of F97 - 2535 were mixed together to give a uniformly

A copying machine commercially available from SANYO ELECTRIC CO., LTD. under the trade name of SFT - Z70 was charged with the obtained developer, and an image was duplicated on a paper for plain paper copy using a test chart No. 1 - R1975 prescribed by THE SOCIETY OF ELECTROGRAPHY OF JAPAN.

Near infrared rays were irradiated to the image - duplicated paper from a dichroic coat type halogen lamp (2 W/cm²) for 2 seconds. Five sheets of the paper and five sheets of non - duplicated paper were put or not at a distance of 5 m from the wall. The distinction was carried out by five men and five women, and point of ten persons was counted, and decolorizing property was evaluated in accordance with the following criteria for evaluation.

[Criteria for evaluation]

- A: 40 to 50 points
- 8: 30 to 39 points
- C: 20 to 29 points
- 0: 0 to 19 points

	3	5	25	20	10	5
			Table 10			
Bxperiment No.		Components of decolorizable toner (parts)	zable toner (part	(6	Physical property of decolorizable toner	ty of
	Resta binder	Near infrared tray-absorbing dye	Decolor1zing agent	Wax	Printed trace after decolorizing	izing
	RB-15 (100)	DY-1 (20)	SE-1 (5)	WA-2 /11		
	RE-15 (100)	DY-1 (20)		WA-2 (1)	٠ ٠	
	RE-16 (100)			(I) # WH	‹ ·	
	RE-16 (100)	DY-3 (2)		ı	۲ ،	
	00	DY-4 (10)		WA-2 (1)	٠,	
	œ	DY-5 (6)		WA-2 (1)	< -	
	RE-19 (100)	DY-6 (2)			۲ ،	
	RE-19 (100)	DY-7 (2)		ì	٠ ،	
	RE-21 (100)	DY-8 (6)			< •	
	RE-21 (100)	DY-1 (1)			< <	
Comparative						
	00-17					
			SE-1 (5)	WA-2 (1)	Q	
		DY-1 (20)	(S) 1-2S	WA-2 (1)		
	RE-22 (100)	DY-2 (2)	SE-2 (6)	; ' ') :	
	RE-23 (100)	DY-3 (2)		ı	י ב	
	RE-17 (100)	DY-4 (10)		WA-2 (1)	ک د د	

From the results shown in Table 10, it can be understood that all of the decolorizable toners containing a light transmittance of not less than 80 % as a resin binder, which were obtained in Experiments 1 to 10, produce little afterimage after decolorizing even though decolorizing time

Example 4

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{Experiments I to IO and Comparative Experiments I to 5]

- 5 Raw malerials shown in Tables 1 to 4 were weighed in a ratio shown in Table 11 so that the total amount could be 800 g. Then, all of the raw materials were thrown into a pressure kneader having an effective volume of 2 t. The kneader was heated, and the kneading temperature of the raw materials was adjusted to 130°C. After the raw materials were kneaded for 15 minutes, the obtained kneaded malerial was taken out from the kneader and cooled to give a solidified product.
- Then, the obtained solidified product was ground with a jet mill and classified by using an air classifier to give toner particles having an average particle diameter of about 12 µm. To the toner particles was added 0.3 % by weight of silica commercially available from NIPPON AEROSIL CO., LTD, under the trade name of R 972 to give a toner.

As a physical property of the toner, decolorizing property was evaluated in the same manner as in Example 2. The results are shown in Table 11.

50	4 0	35	30	20	15	5
			Table II			
Gaperiment		8	Components of decolorizable toner (parts)	olorizable toner	(parts)	
' d	Resin binder	Near infrared ray-absorbing dye	Decolorizing	Wax (polyolefine wax)	Near Infrared ray-reflecting or -absorbing material	Decolorizing property
1	RE-24 (100)	DY-1 (20)	SE-1 (6)	WA-2 (1)	ł	<
N	RE-24 (100)	DY-1 (20)	SP-1 (6)	WA-2 (1)	AD-1 (5)	: <
m	RE-25 (100)	DY-2 (2)		WA-3 (15)	AD-3 (7)	: <
4	RE-25 (100)	DY-3 (2)		WA-3 (5)		: <
¢,	RB-24 (100)	DY-4 (10)	SE-2 (10)	WA-2 (1)	AD-4 (6)	: <
9	RE-24 (100)	DY-5 (20)	SE-1 (10)			: <
2	RE-26 (100)	DY-6 (2)	SE-2 (10)			<
60	RE-25 (100)	DY-7 (2)	SB-3 (10)		AD-2 (10)	: «
σ	RE-24 (100)	DY-8 (10)	SE-2 (20)			: <
01	RE~25 (100)	DY-1 (1)	SE~4 (20)			: <
Comparative Experiment						
**	RE-24 (100)	DY-1 (20)	SE-1 (6)	ı	AD-1 (5)	٥
7	RB-24 (100)	DY-1 (20)		ı		2 (
υn	RE-25 (100)			WA-4 (5)	AD-3 (10)	
*	RE-26 (100)	DY-3 (2)		WA-3 (5)	AD-2 (19)	ם כ
9	RE-24 (100)	DY-4 (10)	SE-2 (10)	WA-2 (0.05)	AP-4 (5)) C

From the results shown in Table 11, it can be understood that all of the decolorizable toners containing a specific polyoletine wax, which were obtained in Experiments 1 to 10 have excellent decolorizing property nevertheless a near infrared ray - reflecting or - absorbing material is contained in the decolorizable toners. Moreover, it can be recognized that the toners obtained in Comparative Experiments 1 to 2 have bad decolorizing property because a polyoletine wax is not contained in the toners, and the toner obtained in

Comparative Experiment 5 has bad decolorizing property because only a little amount of polyolefine wax is contained in the toner. Furthermore, it can be recognized that when a resin having a compatibility with a polyolefine wax is used as a resin binder, as is clear from the result of Comparative Experiment 4, decolorizing property of the toner is lowered because a tens effect is not imparted to the toner by the

Reference Example t

Raw materials shown in Tables 1 to 4 were blended in a blending ratio shown in Experiments 1 to 23 or 0 Comparative Experiment 1 of Table 12 to give a mixture. Then, 20 parts of the mixture and 80 parts of 1.1".2.2" - tetrachloroethylene (hereinafter referred to as "TCE") were blended with stirring to give a uniform

The solution was used as a blank sample in measuring the absorbance of a solution. Components of a blank sample used in each of Experiments 1 to 23 and Comparative Experiment 1 were prepared so as to correspond to the components used in each of Experiments 1 to 23 and Comparative Experiment 1 as

Example 5

20 [Experiments t to 23 and Comparative Experiment t]

Raw materials shown in Tables 1 to 4 were blended in a blending ratio shown in Table 12 to give a mixture. The mixture was kneaded by using a LABO PLAST MILL (model: 20C200, chamber volume 60 mt) commercially available from TOYO SEIKI SEISAKU-SYO., LTD. under the condition that the mixing chamber temperature is 120°C, the filled content in the mixing chamber is 70 % and kneading time is 5 minutes, so that shearing stress was added to the resin binder.

Then, 20 parts of the kneaded material and 80 parts of TCE were mixed together with stirring to give a uniform toner solution.

Absorbance of the toner solution was measured at a wavelength of 640 nm by using a spectrophotometer UV/VIS - 660 commercially available from JASCO CORPORATION. Then, the ratio of absorbance of the toner solution was measured based upon the absorbance of the blank sample obtained in Reference Example 1 of 100. The results are shown in Table 12.

Moreover, discoloration and storage stability of the obtained kneaded material were evaluated in accordance with the following methods. The results are shown in Table 12.

(Discoloration)

The color of the kneaded material was compared with the color of the blank sample with naked eyes, and discoloration was evaluated in accordance with the following criteria for evaluation.

[Criteria for evaluation]

- No change of color is observed in the kneaded material.
- B A slight change of color is observed in the kneaded material, but there is practically no problem.
- A little change of color is observed in the kneaded material, but there is practically no problem.
- D: Oiscolor is obviously observed in the kneaded material, and the kneaded material is not suited for practical uses.

(Storage stability)

After a transparent glass vessel was charged with the kneaded material and preserved for 10 days at morn temperature in an experimental room which could get natural light, the color of the kneaded material was compared with the color of the blank sample with naked eyes, and storage stability was evaluated in accordance with the above ~ mentioned criteria for evaluation of discoloration.

							:	9					
							9000	2					
Experiment	1		Comp	Components of		trable	decolorizable toner (parts)	u-ts)		ļ	Physi decok	Physical properties o decolorizable toner	of
4	_	Resin binder	Near in ray-ab dye	Mear infrared ray-absorbing dye	Decolorizing agent	iring guin	Wax	.,, •	Light fastne stabilizer (Metal oxide)	Light fastness stabilizer (Metal oxide)	Ratio of alsorbance (x)	Discoloration	Storage
~	RB-1	(100)	DY-9	(2)	SE-1	(2.84)	WA-1	19	1	101			
8	RE-1	(300)	OY-1	(3)	SR-1	(2.84)			5	•	20	<	<
ю.	RB-1	(100)	9-XG	(2)	S	(2.84)		3	1 1 4	- `	3	a	Œ
•	<u>-</u> 2	(100)	DY-4	(3)	SE-1	(2.84)	MAN-1	i c	5 6	36	₩ (<	<
۰۵	RE-I	(300)	9740	(3)	SE-1	(2.84)				96	90	<-	<
1 0 1	~G-1	(00)	DY-10	•	SE-1	(2.84)	WA-1-	S G	4-40	•	200	<	<
	RE-1	(002)	DY-7	(5)	SB-1	(2.84)		36	300	36	O P	< 1	<
ro ·	RE-1	(100)	DY-8	(3)	SE-1	2.843			1	٠,	7/	2	C
Ç,	RB-1	(100)	6-XC	(5)		2.84)			5	•	92	<	8
01	RE-1	(100)	DY-1	(5)		(2.84)	/AV-1	36		(0)	80 6	<	<
	- H	(100)	DX-8	(3)	_	(2.84)	•	9	5	(60)	000	< -	<
7:	2 - 3 C	(100)	DY-4	(2)	SE-2 ((3.85)	_	(9)	-40	(60)	7	٠٠	<
7	KE-3	(001)	9	(3)	•	(3.6)	_	2	6-40	(2)	r 0	٠,	< 1
•	1	(100)	DX-10	3	_	2.84)	WA-1	(5)			0 0	< :	:
0	KB-1	(100)	DY-7	(3		(3.5)				(2)	212	5	2
9	RE-I	(007)	DY-8	(3)	SE-3 ((3.5)		(9)			0 0	< -	œ
	-B-1	(100)	6-X0	(20)	-	10)	_	9		(2)	D 10	< -	<
30	RB-1 ((100)	DY-1	(0.2)	_	(20)	_	(3)			0 :	< 1	<
3	_	(100)	DY-6	9	_	(9)	_	(6)	2	(2)	7 0	3 .	2
02	_	300)	- TAG	(01)	_	(20)					0	٠,	<
21	RE-1	100)	DY-5	(0.2)	•	10		19		25	0 0	< (c
22	LE	100)	DY-10	(0.2)							7/	=	ပ
23	RB-3 (100)	DY-7	(00)	2 1-35	(6			1	ĵ.	72	=	~
								•	1	60.	85	<	<
Comparative						•							
	RE-1 (100)	100)	DY-9 ((2)	SE-1 (2	(2.84)	WA-1 (5)	_	1		,	ć	
				.				•			+ 0	2	۵

From the results shown in Table 12, it can be understood that all of the decolorizable toners containing a metal oxide as a light fastness stabilizer obtained in Experiments 1 to 23 do not discolor although the toner obtained in Comparative Experiment 1 shows change of color, and these decolorizable toners have excellent strage stability.

Then, the kneaded materials obtained in Experiments 1 to 23 of Example 5, from which a solvent was removed, were ground by using a jet mill and classified by using an air classifier to give toner particles having an average particle diameter of about 12 µm. To the toner particles was added 0.3 % by weight of silica commercially available from NIPPON AEROSIL CO., LTD. under the trade name of R - 972, and they were mixed by using a Henschel mixer to give a toner

When decolorizing property of the obtained toners was examined by irradiating near infrared rays to the developed image formed from the toners on a copying paper using an aluminum - coated halogen tamp (2 W/cm²), which was placed at a distance of 30 cm from the developed image for one hour, it was observed that all of the toners obtained in each of Experiments had excellent decolorizing property and were suited to practical uses.

[Experiments 24 to 28 and Comparative Experiment 2]

The procedure of Experiment 1 of Example 5 was repeated to give a toner solution except that raw materials shown in Tables 1 to 4 were blended in a ratio shown in Table 13.

The toner solution was coated on a white copying paper with a brush so that a toner layer having a thickess of about 20 to 25 µm in dry could be formed. After the toner layer was dried, near infrared rays were irradiated to the toner layer using a dichroic coal type halogen tamp (2 W/cm²) which was placed at a distance of 30 cm from the paper for 1 hour to decolorize the toner.

Then, the transparency and glossiness of the afterimage of the toner on the paper were evaluated with naked eyes. The results are shown in Table 13.

50	45	35 40	30	25	20	10	5
				Table 13			
Second many		Components	Components of decolorizable toner (parts)	toner (parts)		Physical properties decolorizable toner	ties of
1	Resin binder	Near infrared ray-absorbing dye	Decolorizing agent	Wax	Light fastness stabilizer (Mctal oxide)	Transparency	Glassiness
24	RE-1 (100)	DY-9 (2)	SE-1 (2.84)	WA-1 (5)	04-1 (2)	Translucent	Slightly glossy
26	RE-1 (100)	DY-9 (2)	SE-1 (2.84)	WA-1 (5)	OA-1 (6)	(White) Translucent	Ret
26	RE-1 (100)	DY-9 (2)	SE-1 (2.84)	WA-1 (6)	OA-1 (10)	(White) Translucent	Flat
27	RE-1 (100)	DY-9 (2)	SE-1 (2.84)	WA-1 (6)	OA-1 (50)	(White) Opoque	Fair
88	RE-2 (100)	DY-9 (2)	SE-1 (2.84)	WA-1 (6)	OA-6 (60)	(White) Opaquo (White)	Fist
Comparative Experiment	RE-1 (100)	DY-9 (2)	SE-1 (2.84)	WA-1 (5)	1	Translucent (Light yellow)	Glossy

From the results shown in Table 13, it can be understood that when a metal oxide is contained in the toner as a light lastness stabilizer as shown in Experiments 24 to 28, the gloss of the copyring paper becomes nearly equal to the gloss of an original copyring paper in comparison with the toner not containing a metal oxide as shown in Comparative Experiment 2. Moreover, it can be understood that when the amount of the metal oxide is increased in the loner, the trace of the toner after decolorizing becomes from

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(Experiments 29 to 39 and Comparativ	e Experiments 3 to 41
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Raw materials shown in Tables I to 4 were weighed in a ratio shown in Table I4 so that the total amount could be 800 g, and all of the raw materials were thrown into a pressure kneader having an effective volume of 2 t. Then, the kneader was heated, and kneading temperature of the raw materials was adjusted to 130°C. After the raw materials were kneaded for 15 minutes, an obtained kneaded material was taken out from the kneader and cooled to give a soliditied product.

The obtained solidilied product was pulverized by using a cutter mill to give a loner having a particle diameter of about 1 mm.

As physical properties of the toner, thermal discoloration and light stability were evaluated in accor-dance with the following methods. The results are shown in Table 14.

15 (Thermal discoloration)

Since evaluating the thermal discoloration of the pulverized toner was impossible as it was, the toner was formed into a pellet having a thickness of 2 mm by pressing the toner with a hydraulic press under a pressure of 700 kg-1/cm². As thermal discoloration, a reflection density of the pellet was measured by using a Macbeth densitomer. The higher the value of the reflection density is, the smaller the thermal discoloration is.

(Light stability)

After the pellet was allowed to stand for 48 hours under a fluorescent tamp (illumination intensity: 1500 lux), the reflection density was measured in the same manner as in evaluation of thermal discoloration.

Then, light stability was calculated in accordance with the following equation.

[Light slability (%)]

 = [(Reflection density of the pellet after allowing lo stand)/(Reflection density of the pellet before allowing to stand) x 1001

The higher the value is, the greater the light stability is.

Experiment	1			Сощро	Components of decolorizable toner (parts)	wizable tone	r (parts)		Physical properties of decolorizable toner	rities of
đ	Resin blad	binder	rey ab	Near infrared ray-absorbing dye	Decolorizing agent	Wex	Near infrared roy-reflecting or absorbing material	Light fostness stabiliser (Metal soap)	Thermal discolaration (-)	Light stability (%)
53	P. 12-2	(196)	-¥2		SB-1 (4)	WA-2 (3)	A0-6 (5)	AC-1 (0 3)		
000	RB-2	(100)	DY-2	_	SB-2 (4)		9 QV		- v	77
- 6	F C	(100)	DY-3		SE-3 (4)	WA-2 (3)	AD-6 (5)	MS-2 (1.0)	 	9 6
7 6	666-3	(00)	Ž				AD-6 (5)		4.	
7	7 ((00)	9				AD-5 (6)			• c
* .	KE - 2	(100)	04-10				AD-5 (6)			7 0 6
0 0	H H H	(100)	DY-7	(3)		WA-2 (2)	AD-5 (6)	MS-4 (3.0)		2 0
2 10	KK-3	(100)	9-X0	(3)			AD 6		47.	7 1
37	RB-2	(001)	DY-!	(3)	SE-1 (4)		900			0
38	RB-2	(100)	DY-2	(3)		WA-2 (5)	9		1.40	1.1
39	RB-3	(100)	DY-3	(3)	SE-3 (4)		AD-6 (5)	_	1.33	9 - 2
Comparative				,						
3	RE-3	(100)	DY-7	(2)	SE-3 (4)	WA-2 /2)	3, 3, 4,		,	
•	RE-3 ((100)	DY-8	(5)	SE-4 (6)	WA-2 (3)	(e)	1 1	0.71	30

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From the results shown in Table 14, it can be understood that when a metal soap is used as a light fastness stabilizer in the raw materials for toners, the raw materials are little discolored during kneading the raw materials with heating and have excellent light stability as is clear from Experiments 29 to 39 although the raw materials according to Comparative Experiments 3 and 4 are discolored during kneading.

[Experiments 40 and 41 and Comparative Experiment 5]

Plaw materials shown in Tables I to 4 were weighed in a blending ratio shown in Experiments 43 and 44 or Comparative Experiment 5 of Table I5 so that the total amount could be 5150 to 5200 g. A mixer 5 having an effective volume of 20 t was charged with all of the raw materials, and the raw materials were blended together for 5 minutes while the temperature of the raw materials was adjusted to 25° to 30°C in the mixer. The mixture was taken out from the mixer and a biaxial extruder was charged with the mixture. After the mixture was kneaded, the mixture was cooled to give a solidified product.

The solidified product was formed into a sample having a thickness of 2 mm by pressing the solidified product under a pressure of 700 kg+l/cm² using a hydraulic press.

As physical properties of the sample, image slabilities 1 to 4 were evaluated in accordance with the following methods. The results are shown in Table 15.

First of all, the obtained samples were allowed to stand for 120 minutes under a fluorescent lamp (illumination intensity: 1500 lux).

(A) Image stability 1

The density of color of the sample before allowing to stand and the density of color of the sample after.

allowing to stand are measured by using a Macbeth densitometer. The smaller the absolute value of the difference of density is, the smaller the degree of change of color is.

(B) Image stability 2

The value "L" of the sample before allowing to stand and the value "L" of the sample after allowing to stand are measured by using a Z - E 90 COLOR MEASURING SYSTEM commercially available from NIPPON DENSYOKU KOGYO CO., LTD. The greater the difference is, the greater the degree of change of color is.

(C) Image stability 3

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50

The value "a" of the sample before allowing to stand and the value "a" of the sample after allowing to stand are measured by using a Z - £ 90 COLOR MEASURING SYSTEM commercially available from NIPPON DENSYOKU KOGYO CO., LTD. The smaller the value is, the better the image stability is.

35 (D) Image stability 4

The value "b" of the sample before allowing to stand and the value of "b" the sample after allowing to stand are measured by using a Z-E 90 COLOR MEASURING SYSTEM commercially available from NIPPON DENSYOKU KOGYO CO., LTO. The smaller the value is the better the image stability is.

de 15

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C,

Farmer frame		Comp	onents of	Components of decolorizable toner (parts)	toner (pa	rts)			Physic	pi pro	oertice	of dec	olorizat	Physical properties of decolorizable toner	
đ	Restu binder		Near	Decolorizing	, and	Maht				7	o of It	No. of Image stability	rability		ľ
		[]	absorbing	og ent	<u> </u>	stabilizer.	, i	1		8		3		-	
		ָם ו	dye			(deos	· •	o Afr	120 min.	0 120 min nin	120 nin	o ë	120 mln	o nin	120 mfn
ç	KB-5 (97)		DY-1 (2)	SE-1 (1)	WA-1 (3) MS-1 (1)	MS-1	€	1.22	1.03	33.2	36.9	-11.0	-11.7	1.22 1.03 33.2 36.9 -11.0 -11.1 -15.6 -12.2	-12.2
41	RE-6 (97)		DY-1 (2)	SE-1 (1)	WA-1 (3) MS-1 (0.5)) MS-1	(0.5)	1.07	1.07 0.85	38.3	42.9	-14.9	-14.4	38.3 42.9 -14.9 -14.4 -20.7 -13.6	113.6
Comparative Experiment															
9	RE-5 (97)		DY-1 (2)	SE-1 (1)	WA-1 (3)			0.95	0.61	42.7	54.8	-15.7	-13.6	0.95 0.61 42.7 54.8 -15.7 -13.6 -17.3 -4.3	14.3

From the results shown in Table 15, it can be understood that all of the decolorizable toners containing a metal soap as a light fastness stabilizer obtained in Experiments 40 and 41 have excullent image stability compared with the toner obtained in Comparative Experiment 5.

Example 6

[Experiments 1 to 13 and Comparative Experiments 1 to 3]

Raw malerials shown in Tables 1 to 3, 5 and 6 were blended in a blending ratio shown in Table 18 to give a uniformly blended mixture.

Then, 20 parts of the mixture and 80 parts of 1,1,2,2 - tetrachloroethane were mixed together, and the mixture was sufficiently dissolved to or dispersed in the 1,1,2,2 - tetrachloroethane to give a toner solution.

The toner solution was coaled on a while copying paper with a brush so that a toner layer having a thickness of 20 to 30 µm in dry could be formed, and a solvent was vaporized to remove from the loner layer to give a sample.

As physical properties of the sample, yellowing resistance against fradiation of light and yellowing resistance against heat cycle were evaluated in accordance with the following methods. The results are shown in Table 16.

(Yellowing resistance against irradiation of light)

Near infrared rays were irradiated to the sample from a dichroic coal type halogen lamp (2 W/cm²), which was placed at a distance of 20 cm from the sample, for 1 hour. After decolorizing the sample, a light a having a wavelength of 300 to 400 nm, which was placed at a distance of 48 cm from the sample, was irradiated to the sample through a XENON LAMP FILTER commercially available from ATLAS CO., LTD, for 50, 100 or 500 hours. After that, yellowing of the sample was examined in accordance with JIS - B7754 by using a Canon lade meter. The yellowing resistance against irradiation of light of the sample was evaluated in accordance with the following criteria for evaluation.

[Criteria for evaluation]

- A: No yellowing is observed in the sample when the light is irradiated to the sample for 500 hours.
- B No yellowing is observed in the sample when the light is irradiated to the sample for not less than 100 hours and less than 500 hours.
- C: No yellowing is observed in the sample when the light is irradiated to the sample for not less than 50 hours and less than 100 hours.
- D Yellowing is observed in the sample until the light is irradiated to the sample for 50 hours.

15 [Yellowing resistance against heat cycle]

After a cycle of a procedure comprising irradiating near infrared rays to the sample in the same manner as in the procedure of the yellowing resistance against irradiation of tight, reforming a toner tayer on the sample heating the sample at 80°C for 10 minutes was repeated, yellowing resistance against heat cycle of the sample was evaluated in the same manner as in the above - mentioned yellowing resistance against irradiation of light. The criteria for evaluation is as follows.

[Criteria for evaluation]

- A No yellowing is observed in the sample when the treatment is repeated not less than 50 cycles."
- No yellowing is observed in the sample when the treatment is repeated not less than 20 cycles and less than 50 cycles.
- C. Yellowing is observed in the sample when the treatment is repeated less than 20 cycles.

	5		0	5		0	91 əklət	0	5	0	5
Experiment	Pil			Сощрол	ents of	decolori	Components of decodorizable toner (parts)	r (parts)		Physical decoloriz	Physical properties of decolorizable toner
No	Resto binde	luder.	Near ir ray—abs dye	Mear infrared ray—absorbing dye	Decolo agent	Decolorizing agent	Wox	Ultraviolet absorbing agent	Light fastness stabilizer (Heat- resistant age resistor)	Yellowing resistance against irradiation of fight	Yellowing resistance against heat cycle
-	_	(100)	DY-1	(2)	SE-1	(2.84)	WA-1 (5)) UA-2 (1)	A0-1 (2)	<	<
62	RE-1 C	100)	DY-1	(3)	SE-1	(2.84)	WA-1 (6)	UA-2		: <	: «
P)	ĩ	1001	7	(2)	SE	(2.84)	WA-1 (6)	11A-2		: ∢;	: <:
₹.	7	100)	DY-2	(20)	SB	(9)		UA-2		<	: =
٠	ņ	(00)	DY-3	(10)	- 1 - 2 1	(01)	WA-1 (15)	C-Vn		<	• <
9	Ī	(001)	7	(16)	S	(10)	WA-1 (10	1-VD	AO-4 (26)	<	: <
7	7	(100)	9 2 0	(20)	SE-1	30)	WA-1 (1)	ו-עח		<	: <
∞	RB-2 ()	300)	9-X0	(3)	SB-1	(01)	WA-1 (15)	UA-2		<	: <
Ġ		(001)	DY-7	(3)	- J	(10)	WA-1 (5)	C-43	A2-2 (10)	٠ <	: <
02	_	(300)	DY-8	30	SE-1	(20)	WA-1 (1)	*-Y1		<	: <
1.1	~	(100)	DY-1	Ê	SE-1	(20)			_	: <	٠ <
12	•	(100)	DY-1	(3)	SE-1	(2.84)	WA-1 (6)	UA-2		<	: ==
13	RE-2 (1	6	DY-1	(2)	SE-1	(6.0)	WA-1 (5)	1-Vn	ı	∢	c
Comperative											
-	RE-1 (100)	(00)	DY-1	(2)	SE-1	(2.84)	WA-1 (5)	•	1	c	ď
8		(100)	DY-3	(16)		(10)	_	1	AO-1 (25)	ı U	•
m	RE-3 (1	(100)	DY-1	(10)	SE-1	(01)				U	

From the results shown in Table 16, it can be understood that all of the decolorizable toners containing
an ultraviolet absorbing agent obtained in Experiments 1 to 13 have excellent yellowing resistance against
irradiation of light and yellowing resistance against heat cycle. In addition, when a heat resistant age
resistor is contained in the toner, it can be understood that yellowing resistance is more improved.

Example 7

Experiments 1 to 5]

A resin binder, a near infrared ray - absorbing dye, a decolorizing agent and electrically chargeable fine particles shown in Tables 1 to 3 and 6 were used.

In 75 parts of toluene was dissolved 20 parts of the resin binder, and the near infrared ray - absorbing dye and the decolorizing agent were dissolved therein or mixed therewith in an amount shown in Table 17 based on 100 parts of the resin binder to give a solution. Then, the toluene was removed from the solution dry. After drying, an obtained material was crudely ground, and the material was ground by using a LABO - JET MILL commercially available from NIPPON PNEUMATIC MFG. CO., LTD. to give toner particles having a weight average particle diameter of 10 um. The electrically chargeable fine particles were mixed with the toner particles in a ratio shown in Table 17 based on 100 parts of the toner particles to give from NARA MACHINERY CO., LTD. at a peripheral speed of 80 m/second for 3 minutes so that the a mixture. The mixture was treated by using a HYBRIDIZATION SYSTEM NHS-0 commercially available

As physical properties of the toner, electrically charged amount, printability, decolorizing property and light resistance were evaluated in accordance with the following methods. The results are shown in Table electrically chargeable fine particles could be coaled on the surface of the toner particles to give a toner. 17. (Measuring method for electrically charged amount).

Electrically charged amount was evaluated according to a blow - off method.

2

Aller the toner and a FERRITE CARRIER FB-810 Commercially available from KANTO DENKA charged amount of the mixture was measured when the mixture was blown by using a BLOW - OFF TRIBO CHANGE METER TB - 200 commercially available from TOSHIBA CHEMICAL CO., LTD. under a nitrogen KOGYO CO., LTD. were blended so that the concentration of the toner could be 5 % by weight, the toner and the FERRITE CARRIER were uniformly dispersed, mixed and stirred to give a mixture. The electricatly

gas pressure of 1.0 kg-l/cm² for 30 seconds. 2

(Measuring methods for printability, decolorizing property and light resistance)

A polypropylene bottle (volume: 500 mt) was charged with 5 parts of the toner and 95 parts of a NON-COAT FERRITE CARRIER and they were mixed together by using a rotational system at a speed of rotation of 50 rpm for 30 minutes to give a developer (total weight; 300 g). 8

SUSHITA ELECTRIC CO., LTD, was used and an external high pressure power source was used for a As a printing system, a taser beam printer KX - P4420 commercially available from KYUSYU MAT transfer power source and a developing bias power source, and the printing system was modified so that both a positively charged toner and a negatively charged toner could be used for printing by using the 3

printing system was modified so that the bias voltage of about - 150 V and the transfer voltage of about Originally the above - mentioned printing system is suitable for the negatively charged toner, and a bias voltage of about -550 V and a transfer voltage of about +4000 V can be applied thereto. However, the -4000 V could be applied thereto when the positively charged toner was used. \$

The above - mentioned bottle having the developer was lixed in the above - mentioned printing system in a usual method and 1000 sheets of paper were printed in accordance with a regular printing method.

(i) Printability Ş

After 1000 sheets of paper were printed as mentioned above, the surface of the 1000th printed paper was observed with naked eyes under natural light. The state of printing was examined and evaluated in accordance with the following criteria for evaluation.

(Criteria for evaluation)

S

- Colorability of printing is excellent and no fogging is observed in the printed paper. ₹ # Ö

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Colorability of printing is good but logging is observed a little in the printed paper. Colorability of printing is bad and logging is observed much in the printed paper.

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(ii) Decolorizing property

After half of the 1000th printed paper was covered with an aluminum film and was laid on a hot plate at 70°C, the printed paper was allowed to stand for 3 minutes. Then, near infrared rays were irradiated to the printed paper for 8 seconds from an aluminum - coated halogen lamp for irradiation of near intrared rays (2 W/cm²) which was placed at a distance of 30 cm from the printed paper, and the printed paper was observed with naked eyes and decolorizing property was evaluated in accordance with the following criteria for evaluation. 'n

[Criteria for evaluation] 5

- The toner is completely decolorized. ≆ ä ö
- The color of the toner is slightly remained in the printed paper.
- The toner is a little decolorized, but the remained color of the toner is obviously observed in the printed paper.

(iii) Light resistance

5

In the above -mentioned procedure for producing a printed paper, in order that toner particles were uniformly thinly extended over the paper and near infrared rays were uniformly and sufficiently irradiated, a non - fixed printed paper was produced. Near intrared rays were irradiated to the printed paper for 60 seconds from an aluminum - coated halogen lamp for irradiation of near infrared rays (0.2 W/cm²), which was placed at a distance of 30 cm from the printed paper. 2

Then, the non-fixed printed paper to which near infrared rays were uradiated and the non-fixed printed paper to which near infrared rays were not irradiated were introduced into a fixing system to fix the printed matters. Difference of colorability of printed matters after fixing between the above - mentioned two non - lixed printed papers was observed with naked eyes, and evaluated in accordance with the followig criteria for evaluation. 2

[Criteria for evaluation] 8

- No difference of colorability is observed between the two. **₹ 66 ∵**
- A little difference of colorability is observed between the two.
- Much difference of colorability is observed between the two.

[Experiments 6 to 10]

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A resin binder, a near infrared ray - absorbing dye, a decolorizing agent and an electrically chargoable fine particles shown in Tables 1 to 3 and 6 were used.

The near infrared ray - absorbing dye and the decolorizing agent were added to the resin binder in a ratio shown in Table 17 based on 100 parts of the resin binder and they were dissolved in 200 parts of methylene chloride to give a solution. Then, the methylene chloride was removed from the solution. After an obtained material was coasely pulverized, the material was pulverized by using a LABO-JET MILL commercially available from NIPPON PNEUMATIC MFG. CO., LTD. to give toner particles having a weight average particle diameter of 10 um. The, electrically chargeable fine particles were mixed with the toner particles in a ratio shown in Table 17 based on 100 parts of the toner particles, and the electrically chargeable line particles were coated on the surface of the toner particles in the same manner as in Experiment 1 of Example 7 to give a toner. \$ ŝ

Physical properties of the toner were evaluated in the same manner as in Experiment 1 of Example 7, The results are shown in Table 17.

Experiment 11]

8

A resin binder, a near intrared ray - absorbing dye, a decolorizing agent, an electrically chargeable line particles and an electric charge regulator shown in Tables 1 to 3, 5 and 6 were used. 55

In 200 parts of methylene chloride were dissolved 100 parts of the resin binder, 2.1 parts of the near intrared ray - absorbing dye. I part of the decolorizing agent and 2.7 parts of the electric charge regulator. and they were kneaded to give a solution. Then, the methylene chlorids was removed from the solution.

After an obtained material was coasely pulverized, the material was pulverized by using a LABO - JET MILL commercially available from NIPPON PNEUMATIC MFG. CO., LTD to give loner particles having a weight average particle diameter of 10 µm. With 100 parts of the toner particles were mixed 20 parts of the electrically chargeable line particles, and the electrically chargeable line particles were coated on the 5 surface of the toner particles in the same manner as in Experiment 1 of Example 7 to give a toner.

Physical properties of the toner were evaluated in the same manner as in Experiment 1 of Example 7. The results are shown in Table 17.

[Comparative Experiment 1]

The procedure of Experiment 1 of Example 7 was repeated to give a toner except that the electrically chargeable line particles were not coated on the surface of the toner particles.

Physical properties of the loner were evaluated in the same manner as in Experiment 1 of Example 7. The results are shown in Table 17.

(Comparative Experiment 2)

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The procedure of Experiment 11 of Example 7 was repeated to give a loner except that the electrically chargeable fine particles were not coaled on the surface of the toner particles.

Physical properties of the toner were evaluated in the same manner as in Experiment 11 of Example 7. The results are shown in Table 17.

> 2012 bias word den 4

.(4)) 4% PS 67.

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 $\mathfrak{m}!=\mathfrak{q}_{\mathfrak{p}}$

50	45		40	35	30	25	20	15	10	5
					Teby	<u>ТаМе 12</u>				
xperiment			Components	of decolorizable toner (parts)	toner (part	(8)	Physical	properties	Physical properties of decolorizable toner	toner
4	Resin binder	binder	Near infrared rey- absortsing dye	Decolorizing agent	Electric charge regulator	Mectalcally chargeable fine porticles	Electrically charged amount (# C/g)	Print- ability	Decolorizing property	Light
	RB-24			(5) :-88		78-1 (20)	-22.0			
N (RE-24	-			ı		-21.4	: <	٠.	٠.
η,	HE-24	100	_		1		+18.1	: <	: <	٠.
• 4	7	(100)			1	TTE-3 (10)	-17.6	: <	: <	٠ <
n 4	1010-24	(100)		Sg-6 (10)	ı	TE-3 (10)	-18.6		: <	٠.
.	KE-Z4	(001)		_	ı	TE-2 (10)	+17.4	<	: <	٠.
• •	K-G-24	000			ı	TE-3 (15)	-13.3	<	: <	٠ <
	12-87	(100)			1	TE-4 (10)	-21.3	٠ <	: <	: •
, ·	RE-24	000			1	TE-6 (10)	6.91-	: <	€ <	٠.
2 :	RE-25	000		(9)	,	12.6 (10)	-18.3	: <	: «	٠,
11	RE-26	(100	DY-1 (2.1)	E 1-35	CR-2 (2.7)		-22.8	: ∢	< <	٠,
Comparative Experiment										
-	RB-24 (100)	(100)	DY-1 (6)	SB-1 (6)	ı	1	* * * +	c	•	(
N	RB-26 (100)	(100)	DY-1 (2.1)		CR-2 (2.7)	1	-2.0	ט נ	< <	ں د

From the results shown in Table 17, it is understood that all of the decolorizable toners obtained in Experiments 1 to 11 of Example 7 have excellent tribo - electrically charged amount, excellent printability. 55 excellent decolorizing property and excellent light resistance.

Example 8

In the following Experiments and Comparative Experiments, raw materials shown in Tables 1 to 3 and 5 were used.

(Experiment 1)

Atter 20 parts of a rest binder (RE - 24) was dissolved in 75 parts of toluene, 2 parts of a near infrared ray - absorbing dye (DY - 2) and 2 parts of a decolorizing agent based on 100 parts of the resin binder were dissolved therein and mixed therewith. A solvent was vaporized therefrom to dry an obtained product. The dred product was coasely pulverized, and further pulverized with a jet mill commercially available from 1.5 parts based on 100 parts of the toner with the toner. The mixture was treated for 3 minutes at a NIPPON PNEUMATIC MFG. CO., LTD., and a toner having a weight average particle diameter of 11 µm was obtained. An electric charge regulator having postively chargeable property was mixed in an amount of peripheral speed of 80 m/second by using a HYBRIDIZATION SYSTEM commercially available from NARA MACHINERY CO., LTD, to cover the surface of the toner particles with the electric charge regulator. The obtained toner showed a blue color. 6

As physical properties of the obtained toner, tribo-electrically charged amount and decolorizing properly were evaluated in accordance with the following methods. The results are shown in Table 18.

(Tribo - electrically charged amount)

2

In accordance with a blow - off method, tribo - electrically charged amount was mesured in the same manner as in Example 7.

(Decolorizing property)

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The amount 4 parts of the toner was mixed with 100 parts of a silicone resin-coated carrier commercially available from POWDERTECH CO. LTD. under the trade name of F97 - 2535 to give a uniformly blended developer. 8

The obtained developer was provided in a copying machine commercially available from SANYO ELECTRIC CO., LTD. under the trade name of SFT - 270, and a reproduction of an image was carried out using a test chart No. 1 - R1975 prescribed by THE SOCIETY OF ELECTROGRAPHY OF JAPAN on a paper for plain paper copy. 35

source of near infrared rays, the paper was adhered to a vertical walt, distinguishing was carried out at a After the obtained paper was irradiated for 3 seconds with an aluminum - coated halogen lamp as a distance from 5 m from the wall with naked eyes. Evaluation of decolorizing property was carried out in accordance with the following criteria for evaluation.

[Criteria for evaluation] \$

- Color of the toner is completely disappeared.
 - Color of the toner is remained slightly.
- Color of the toner is disappeared a tittle, but the existence of the toner is obviously observed. ≆ ä ö ö 5
 - Decolorizing of the toner is not observed.

[Experiment 2]

The procedure of Experiment 1 of Example 8 was carried out except that DY - 3 as a near infrared ray - absorbing dye and SE - 2 as a decolorizing agent were used to give CR - 1 - coated toner particles. The obtained toner showed a blue color. 8

As the physical properties of the obtained toner, tribo - electrically charged amount and decolorizing property were examined in the same manner as in Experiment 1 of Example 8. The results are shown in

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Experiment 3]

average particle diameter of 11 μm. After 2.0 parts of an electric charge regulator (CR = 2) was mixed with The procedure in Experiment 1 of Example 8 was carried out to give toner particles having a weight 100 parts of the toner particles, and the surface of the toner particles was covered with CA - 2 in the same manner as in Experiment 1 of Example 8.

As the physical properties of the obtained toner, tribo - electrically charged amount and decolorizing property were evaluated in the same manner as in Experiment 1 of Example 8. The results are shown in Table 18.

[Experiment 4]

6

The amount 100 parts of a resin binder (RE - 24). 2 parts of a near infrared ray - absorbing dye (DY - 2) and 2 parts of a decolorizing agent (SE-1) were mixed with 200 parts of methylone chloride, and the mixture was knoaded. Then, the methylene chloride was removed from the mixture. After the product was coasely puiverized, the product was pulverized with a jet inill to give toner particles having a weight average particle diameter of 11 µm. 2

The amount 100 parts of the toner particles and 3 parts of CR - 3 were mixed together, and the mixture was treated for 3 minutes at a peripheral speed of 80 m/second by using a HYBRIDIZATION SYSTEM commercially available from NARA MACHINERY CO., LTD. to cover the surface of the toner particles with CR = 3. The obtained toner showed a blue color. 8

As the physical properties of the obtained toner, tribo - electrically charged amount and decolorizing properly were evaluated in the same manner as in Experiment 1 of Example 8. The results are shown in Fable 18.

(Experiment 5)

23

To 100 parts of toner particles having a weight average particle diameter of 11 um propared in the same manner as in Experiment 4 of Example 8, a solution prepared by dispersing 1.5 parts of an electric charge regulator in 20 parts of methanol was added and stirred. The mixture was dried in a FLOW COATER MULT - PURPOSE commercially available from NARA MACHINERY CO., LTD, at a temperature of the inside of apparatus of 55°C for 30 minutes, to homogeneously cover the surface of the toner particles with CR - 4 to give a toner. The obtained toner showed a blue color,

8

As the physical properties of the obtained toner, tribo - electrically charged amount and decolorizing property were evaluated in the same manner as in Experiment 1 of Example 8. The results are shown in Table 18 3

(Experiments 6 to 18)

binder, a near infrared ray - absorbing dye, a decolorizing agent and an electric charge regulator shown in The procedure of Experiment 1 of Example 8 was carried out except, that as raw materials, a resin Tables 1 to 3 and 5 were used in a ratio shown in Table 18. The surface of the toner was covered with the electric charge regulator in the same manner as in Experiment 1 of Example 8. \$

As the physical properties of the obtained toner, tribo - electrically charged amount and decolorizing property were evaluated in the same manner as in Experiment 1 of Example 8. The results are shown in ţ

[Comparative Experiment 1]

Atter 20 parts of a resin binder (RE - 24) was dissolved in 75 parts of toluene, 2 parts of a near infrared ray absorbing dye (DY - 2), 2 parts of a decolorizing agent (SE - 1) and 1.5 parts of an electric charge regulator having a positively chargeable property were dissolved in 100 parts of the resin binder, and mixed therewith so that the mixture was homogeneously mixed. Then, the solvent was removed from the solution, and the obtained product was dried. After the dried product was coasely pulverized, the product was further pulverized with a jet mill to give a toner having a weight average particle diameter of 11 um in which the electric charge regulator (CR - 1) was contained. The obtained toner showed a blue color 8 22

property were evaluated in the same manner as in Experiment 1 of Example 8. The results are shown in As the physical properties of the obtained foner, tribo - electrically charged amount and discolorizing

Table 18.

The copied image had mush logging, and was not a suitable one.

(Comparative Experiment 2)

With 100 parts of a resin binder, 2 parts of a near infrared ray absorbing dye (DY - 2), 2 parts of a decolorizing agent (SE - 2) and 1.5 parts of an electric charge regulator, 200 parts of methylene chloride was mixed. After the mixture was molten and kneaded, methylene chloride was removed from the mixture. The product was coasely pulverized and then pulverized with a jet mill to give a toner having a weight average particle diameter of 11 µm in which the electric charge regulator was contained. The obtained toner showed a blue color.

As the physical properties of the obtained loner, tribo - electrically charged amount and decolorizing property were evaluated in the same manner as in Experiment 1 of Example 8. The results are shown in Table 18.

The copied image had much logging, and was not a suitable one.

[Comparative Experiments 3 to 6]

The procedure of Experiment 1 of Example 8 was carried out except that as raw materials, a resin binder, a near infrared ray - absorbing dye a decolorizing agent and an electric charge regulator shown in Tables 1 to 3 and 5 were used in a ratio shown in Table 18. A toner in which an electric charge regulator was contained was obtained in the same manner as in Comparative Experiment 1 of Example 8.

As the physical properties of the obtained loner, tribo-electrically charged amount and decolorizing property were evaluated in the same manner as in Experiment 1 of Example 8. The results are shown in

The copied image had much logging, and was not a suitable one.

	s	5	00	25	!5 20	5
			Table 18	89		
Experiment	ပို	Components of decol	decolorizable toner (p	(perts)	Physical properties	of decolorizable con
	Resin binder	Near infrared ray-absorbing dye	Decolorizing agent	Rectric charge reculator	dectrically amount	Decolorizing proporty
	١.	L				
	_	DY-2 (2)	SE-1 (2)	CR-1 (1.5)	0 66+	
	KE-24 (100)	DY-3 (2)		(S 1) 1-80	2000	<
	_	-				<
	Ţ	_			97.1-	<
	RB-24 (100)				-22.3	<
	RE-24 (100)	•		(K-4 (1.6)	+26.3	<
		100			+30.8	: <
	^				+21.6	
	•		(2) 2-575	_	+19.8	٠.
	001	02. 0. (2)	_		-17.0	٠.
	•	DY-2 (Z)	SB-1 (2)	CR-3 (2.6)	-24.3	٠.
	DB-26 (100)			-	+25.5	۲.
	~			_	2000+	< •
	(100) DE-24 (100)				+	۲,
	•	_		-	9 00 00 00 00 00 00 00 00 00 00 00 00 00	< -
	_ `	DX-6 (2)	SE-2 (2)	CR-2 (2.0)	0.61-1	۲.
	NE-20 (100)		_	-	0.00	< -
	=;		SE-3 (2)	_	7 7 7	< ∙
	HE-26 (100)	DY-8 (2)	•	CR-6 (2.6)	+29.8	< •
Comparative						
TX DG. I ment						
	RE-24 (100)		SR-1 (2)	08-1-80	- 91	1
	Ç					۵
	C			(1.0)	45.2	۵
					-3.8	c
	:				14.1	16
	(001)	DI-2 (2)	(2) 1-35		+7.6	ه د
				CR-5 (2.5)	+9.2	3 (

From the results shown in Table 18, all of the toners obtained in Experiments 1 to 18 have excellent so electrically charging property and excellent decolorizing property.

Example 9

(Experiments 1 to 10)

was charged with a resin binder A, a near infrared ray-absorbing dye, a decolorizing agent and a wax in a ratio shown in Table 19. After the mixture was heated to a temperature shown in Table 19 and kneaded so that the mixture was homogeneously dispersed, an obtained product was cooled to room temperature to give a kneaded material.

about 5 mm, a pressure kneader was charged with the pulverized material together with a resin binder B After the kneaded material was pulverized with a hammer mill so that the particle diameter was at most having an average particle diameter of at most about 10 mm as shown in Table 1. When the temperature was attained to the temperature shown in Table 19, the mixture was kneaded with heating at that temperature for the time shown in Table 19. After the product was taken out from the pressure kneader and cooled to room temperature, the cooled product was pulverized with a hammer mill to give a decolorizable õ 2

As the physical properties of the obtained toner, colorability, decolorizing property, dispersibility, concentration of image and offset resistance were evaluated in accordance with the following methods. The results are shown in Table 19.

(A) Cotorability 2

The toner was placed on the sample table, the surface color was measured with a Z - E 90 COLOR MEASURING SYSTEM commercially available from NIPPON DENSYOKU KOGYO CO., LTD., and the colorability was evaluated in accordance with the following criteria for evaluation. In the values of L. a. b.

The smaller the value of "b" is, the greater the blue color becomes. The greater the value "b" is, the color elements, the value of "b" was adopted as the color of the decolorizable toner.

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greater the yellow color becomes. When the value of "b" is zero, the color is gray,

Criteria for evaluation}

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- The value of "b" is not more than 10.
- The value of "b" is more than 10 and not more than 3.
 - The value of "b" is more than 3.

(B) Decolorizing property 2

The amount 4 parts of the toner was blended with 100 parts of a silicone resin-coated carrier commercially available from POWDERTECH CO. LTD under the trade name of F97 - 2535, and they were uniformly dispersed to give a developer.

ELECTRIC CO. LTD. and the reproduction of an image was carried out using a test chart No. 1-R1975 prescribed by THE SOCIETY OF ELECTROGRAPHY OF JAPAN on a paper for plain paper copy. The obtained developer was set in a copying machine SFT - 270 commercially available from SANYO ş

After near intrared rays were irradiated to the obtained paper at a distance of 30 cm from the paper for 5 seconds with an aluminum - coated halogen lamp, decolorizing property was evaluated with naked eyes in accordance with the following criteria for evaluation. ç

(Criteria for evaluation)

- Color is completely disappeared, and yellowing is not generated. ≆ ä ö 3
- It takes at least 5 seconds for decolorizing the toner, but yellowing is not occurred.

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Yellowing is observed.

(C) dispersibility

The toner was coasely pulverized with a hammer mill so that its particle diameter is at most 5 mm or so and bonded with an epoxy resin. The product was sliced with a microtome to give a film, and the part of the decolorizable toner was observed by means of a microscope under visible light. The dispersibility was evaluated in accordance with the following criteria for evaluation. 22

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[Criteria for evaluation]

- Both the near infrared ray absorbing dye and the decolorizing agent are not coagurated Agglomerate is observed, but the agglomerate of at least 1 µm is not existed ₩ ₩
 - Agglomerate is observed, but the agglomerate of at least 3 µm is not existed. ö ت
 - Agglomerate of more than 3 µm is existed.

(D) Concentration of image

Concentration of image was measured using a colored filter by means of a Macbeth reflection densitometer and evaluated in a ccordance with the following criteria for evaluation. 5

[Criteria for evaluation]

The measured value is at least 0.9. æ æ ö ö

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- The measured value is at least 0.8 and less than 0.9.
 - The measured value is at least 0.6 and less than 0.8.
 - The measured value is less than 0.6.

(E) Offset resistance 8

Twenty sheets of paper were reproduced by means of a copying machine commercially available from SANYO ELECTRIC CO., LTD. in which a silicone oil-impregnated cleaning pad of a lixing roller was removed, and the number of ottset -generated papers was counted and evaluated in accordance with the following criteria for evaluation. 22

[Criteria for evaluation]

- Number of offset generated papers is zero. ₹ 60
- Number of offset generated papers is 1 or 2.

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- Number of offset generated papers is 3 or 4. ö
- Number of offset generated papers is at least 5. ä

[Comparative Experiments 1 and 2]

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A pressure kneader was charged with a resin binder B, a near infrared ray - absorbing dye and a decolorizing agent without a resin binder A, and they were kneaded in conditions shown in Table 19 Atter the mixture was heated and kneaded for 5 minutes (Comparative Experiment 1) or 1 minute (Comparative Experiment 2) at a temperature shown in Table 19, the mixture was cooled to room temperature, and a product was obtained.

was pulverized with a hammer mill to give a decolorizable toner having an average particle diameter of about 12 µm. The obtained product

The physical properties of the obtained toner was evaluated in the same manner as in Experiments 1 to 10 of Example 9. The results are shown in Table 19.

Table 19

Experiment		onditions for pre	paring molten k	neaded mater	ial	Conditions f	or preparing	
No.	Resin binder (parts)	Near Infrared-ray absorbing dys (parts)	Decolorizing agent (parta)	Wox (parts)	Kneading temperature (°C')	Resin binder B (parts)	Heating compensature (°C')	ficating time (min)
į.	REA-1 (20)	DY-1 (2)	SE-1 (2)	WA-1 (3)	110	REB-1 (80)	130	1
2	REA-2 (20)	DY-1 (2)	SFI-1 (2)	WA-1 (3)	110	REB-1 (80)	130	1
3	REA-3 (20)	DY~1 (2)	SE-1 (2)	WA-1 (3)	110	REB-1 (80)	130	1
4	REA-1 (20)	DY-2 (10)	SE-2 (10)	-	110	REB-2 (80)	130	ì
5	REA-2 (20)	DY-3 (20)	SB-2 (20)	-	110	REB-2 (80)	130	ì
6	REA-3 (20)	DY-4 (20)	SE-2 (20)	-	110	REB-2 (80)	130	ì
7	REA-1 (30)	DY-6 (2))	SE-4 (2)	WA-1 (3)	110	REB-3 (70)	130	1
8	REA-2 (40)	DY-6 (20)	SE-4 (20)	WA-1 (3)	110	REB-3 (60)	130	1
9	REA-3 (50)	DY-7 (2)	SB-4 (2)	WA-1 (3)	110	REB-3 (50)	130	1
10	REA-4 (60)	DY-8 (2)	SB-1 (2)	WA-1 (3)	110	RPB-1 (40)	130	ì
Comparative Experiment		-		· · · ·		· · · · · ·		
1	REA-1 (100)	DY-1 (2)	SR-1 (2)	₩A-1 (3)	130	_	-	-
2	REA-2 (100)	DY-1 (2)	SB-1 (2)	WA-1 (3)	110	-	-	-

- continued -

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		Physical pr	operties of deco	orizadio toner	
Experiment No.	Colorability	Decolorizing property	Dispersibility	Concentration of image	Offset resistance
1	Λ	Λ	Α.	٨	A
•	A	A	A	A	Λ
2	. <u>.</u> A	Ä	A	٨	A
3	Δ.	Ā	A	Λ	A
4	A	Λ	A	A	٨
5	Λ.	Ä	٨	A	٨
6	, A	Α.	Ā	٨	٨
7	· -	,, ,	A	٨	٨
8	٨	Λ Λ	Ā	A	٨
9	٨	Å	Λ	A	Λ
10	A				
Comparative Experiment				n	٨
1	С	С	Λ.		D
2	Α,	В	٨	Λ	

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å A first solution and a second solution, were prepared by mixing the raw materials shown in Tables 1, 2 and 4 to θ in a mixing ratio shown in Table 20. From the results shown in Table 19, it can be understood that a decolorizable toner having excellent colorability, excellent decolorizing property, excellent dispersibility, excellent concentration of image and excellent offset resistance at a time can be obtained in accordance with the above method. [Preparation Example 1 of Master Batch (Preparation No. MB - 1 to 8)]

After the first solution and the second solution were mixed together, an organic solvent was removed from the solution under a reduced pressure to give a mass.

The mass was then pulverized for 50 hours by using a ball mill to give a master batch.

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(20) (60) (60) (60)

AO-6 (3)

(2)

(0.6)(2)

AO-7

solution (parts)

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Lents

Light fastness stabilizer second

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Preparation No.		Components of master batch (parts)
	Resin binder	Near infrared ray - absorbing dye	Light fasiness stabilizer
MB-9	RE - 1 (10)	DY - 1 (2)	AO - 7 (2)
M8 - 10	RE - 6 (10)	DY - 2 (2)	-
MB - 11	RE - 7 (10)	DY - 1 (2)	-
MB - 12	RE -8 (10)	DY - 3 (2)	-
MB - 13	RE - 1 (10)	DY - 1 (2)	AO - 6 (2)
MB - 14	RE -5 (10)	DY - 2 (2)	AO - 6 (2)
MB - 15	RE - 5 (10)	OY - 4 (2)	AO - 11 (1)
MB - 16	RE - 7 (10)	OY - 1 (2)	-
MB - 17	RE - 1 (10)	DY - 3 (2)	AO - 7 (3)

Example 10

20 [Experiments I to 20]

In accordance with the adding method of a master batch shown in Table 22, the raw materials shown in Tables 1, 3 and 4 and the master batch obtained in Preparation Nos. MB - 1 to 17 were heated to melt and kneaded at a temperature of 130°C with a kneader shown in Table 22. In Table 22, EX denotes a braxial 25 kneading extruder and KN denotes a pressure kneader. After that, the kneaded molten material was cooled to give a mass and the mass was pulvelized with a culter mill and a jet mill to give a pulvelized inaterial The pulvelized material was classified with a wind - force classifier to give a toner having a particle charneter of 5 to 20 µm.

The meanings of the adding methods of a master batch 1 to 4 are as follows.

(Adding method of a master batch 1)

After the master hatch and the component for decolonizing were uniformly mixed together with a kneader, the mixture was heated to melt and kneaded with a biaxial kneader or a pressure kneader,

(Adding method of a master batch 2)

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The master batch was introduced into the component for decolorizing while kneading and heating to melt the component for decolorizing with a biaxial kneader so that they were mixed in a prescribed ratio.

(Adding method of a master batch 3)

The component for decolorizing was heated to melt and kneaded for 10 minutes, and the master batch was added thereto. Then, they here heated to melt and kneaded.

(Adding method of a master batch 4)

The component for decolorizing was healed to melt and kneaded with a lirst biaxial kneader. The master batch was heated to mell and kneaded with a second biaxial kneader.

The mollen master batch was introduced into the first braxial kneader so that they were mixed in a prescribed ratio.

[Preparation Example 2 of Master Batch (Preparation No. MB - 9 to 17)]

The raw malerials shown in Tables 1, 2 and 5 were blended logether in a blending ratio shown in Table 21 and healed to melt and kneaded at a temperature of 130 °C by using a biaxial extruder commercially available from IKEGAI CORP, under the trade name of PCM - 30.

As to Preparation Nos. MB - 16 and 17, the obtained kneaded mollen malerial was used as a master 55 batch as it was.

As to Preparation Nos. MB - 9 to 15, the obtained kneaded molten material was cooled to give a mass, and the mass was pulverized for 50 hours with a ball mill to give a master batch.

Table 22

Experiment		Component	s of decoloriza	ible toner (parts)		
Na.			for decolorizing			Method for
	Resin binder	Decolorizing agent	Yax	Near infrared ray-reflecting or -absorbing material	Master batch	adding master batch
1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18	RE-1 (90) RE-5 (90) RE-5 (90) RE-5 (90) RE-5 (90) RE-6 (90)	SE-1 (3) SE-2 (5) SE-1 (6) SE-1 (2) SE-1 (1) SE-2 (0.6) SE-2 (3) SE-4 (4) SE-4 (2) SE-1 (3) SE-4 (2) SE-1 (5) SE-1 (5) SE-1 (2) SE-1 (4) SE-1 (1) SE-2 (6) SE-1 (1) SE-3 (1) SE-3 (4)	WA-1 (3) WA-1 (3) WA-1 (5) WA-1 (3) WA-1 (4) WA-1 (3) WA-1 (5) WA-1 (3) WA-1 (2) - WA-1 (3)	AD-6 (5) AD-5 (6) AD-5 (6) AD-5 (6) AD-6 (6) AD-6 (5) AD-6 (5) AD-5 (6) AD-5 (6) AD-5 (6) AD-6 (5) AD-6 (5) AD-6 (5) AD-6 (5) AD-6 (5) AD-5 (6) AD-5 (6) AD-5 (6) AD-5 (6) AD-5 (6) AD-6 (5) AD-6 (5) AD-6 (5) AD-6 (5) AD-6 (5)	MB-1 (10) MB-2 (7) MB-3 (10) MB-4 (7) MB-5 (10) MB-6 (10) MB-7 (5) MB-9 (10) MB-10 (10) MB-11 (6) MB-12 (10) MB-13 (10) MB-14 (10) MB-16 (10) MB-16 (10) MB-17 (10) MB-17 (10) MB-17 (10) MB-17 (10) MB-17 (10) MB-16 (10)	1 1 1 2 2 2 3 3 1 1 1 1 2 2 2 3 3 3 1

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Experiment	K	neader	Physical prop	erties of decoid	orizable toner
No.	Pirst process	Second process	Discoloration resistance	Light stability	Decolorizing property
1	EX	-	٨	Λ	
2	ТX	_	· ·		A
3	KN	-	A	Λ.	A.
4	KN	_	A .	Λ.	٨
5	EX	-	Λ.	Λ	A
6	EX		Δ.	٨	Λ
7	KN		Λ.	A.	٨
8	KON	_	Ą	٨	A
ă	EX	-	٨	٨	٨
10		-	Λ	٨	A
11	KN	-	Λ	٨	Ā
	EX	-	٨	A	 A
12	KN	-	Λ	Ā	Ä
13	EX	-	A	 A	A.
14	EX	-	Λ.	, , , , , , , , , , , , , , , , , , ,	Λ.
15	KN	-	Ä	A.	Λ.
16	KN	-	Å	Λ. •	Λ.
17	KN		A A	Λ.	Λ.
18	EX	EX	Λ Α	Λ	A
19	EX	EX	<i>N</i> .	A	٨
20	ĒΧ		Λ.	Ą	٨
	υΛ .	EX	A	A ·	٨

The mixture was then blended with a carrier commercially available from POWDERTECH CO., LTD. under the trade name of F883 - 1025 so that the content of the toner was 7 % by weight to give a twocomponent type developer To 100 parts of the obtained toner, 0.1 part of finely powdered silica commercially available from NIPPON AEROSIL CO., LTD, under the trade name of Aerosil R -972 was added, and they were blended to

obtained two - component type developer by means of an electrostatic reproduction machine commercially available from RICOH COMPANY, LTD under the trade name of FT - 4525. After that, the reproduction print that of the previously printed image, find the procedure was reciprocated so that the difference became superposed on the image - printed portion. The Macbeth density of the printed image was compared with was set in a tray for paper, and the reproduction was again carried out so that the printed image was A black solid was used as a manuscript and a reproduction of the black solid was carried out with the

As the physical properties of the toner, discoloration resistance, light stability and decolorizing property of the sample were examined in accordance with the following methods. The results are shown in Table 22, :005 Whon the difference of the Macbeth density was ±0.05, the printed paper was used as a sample.

(Discoloration resistance)

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The toner was formed into a plate having a thickness of about 2 mm and a diameter of about 50 mm by using an oil press, and the reflection density of the plate was measured with a Macbeth densitometer for 20

The average was measured and the discoloration resistance was evaluated in accordance with the following criteria for evaluation.

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23

[Criteria for evaluation]

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Reflection density is at least 1.10. ë ö

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- Rellection density is at least 0.91 and less than 1.10.
- Reflection density is at least 0.71 and less than 0.91.
 - Reflection density is less than 0.71.

(Light stability) 2

The reflection density A of the sample was measured with a Macbeth densitometer. After the sample was allowed to stand for 24 hours under a fluoroescent lamp (illumination intensity: 1500 tux), the reflection density B was measured in the same manner as the above. The light stability was calculated in accordance with the following equation. S.

[Light stability]

= (Reflection density B)-(Reflection

density A) x 100

2

The light stabuity was evaluated in accordance with the following criteria for evaluation.

(Criteria for evaluation)

Reflection density is at least 80 %. ¥ a ∵ ö

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- Reflection density is at least 61 % and less than 80 %. Reflection density is at least 41 % and less than 61 %.
- Reflection density is less than 41 %.

(Decolorizing property) \$

The sample was allowed to stand in a thermostatic chamber of 60°C while irradiating a light with an incomment of 10 cm from the sample. The sample was chaeved with aluminum - coated halogen lamp at a distance of 10 cm from the sample. The sample was observed with naked eyes from the window of the chamber, and the time when the color of the toner was disappeared was ÷

The average decolorizing property was evaluated in accordance with the following criteria for evaluation.

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[Criteria for evaluation]

- Decolorizing time is less than 10 seconds. 8
- Decolorizing time is at least 10 seconds and less than 30 seconds.
- Decolorizing time is at least 60 seconds or the color is not disappeared. Decolorizing time is at least 30 seconds and less than 60 seconds.
 - [Comparative Experiments 1 to 16]

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The raw materials shown in Tables 1 to 5 were mixed together in a blending ratio shown in Table 23 with a kneader and heated to melt and kneaded at a temperature of 130°C. In Table 23, EX denotes

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braxial kneading extruder and KN denotes a pressure kneaderr. After that, the kneaded minition material was cooled to give a mass and the mass was pulvelized with a cutter init and a jet mill to give a pulvelized material. The pulvolized material was then classified with a wind -force classifier to give a toner having a particle diameter of 5 to 20 µm. The physical properties of the toner were examined in the same manner as in Experiments 1 to 20 of Example 10. The results are shown in Table 23.

Comparative			3	apone	ints of	decot	orizat	de to	Components of decolorizable coner (parts)	ıts)		i	2		Physical properties decolorizable toner	ties of
Experiment No.	Resin binder		Near infra ray etbe	Near infrared rny- absorbing dye	Decolorizing agent	rizing	₩ A		Light featness stabilizer	50	Mear Infrared ray- reflecting or absorbing material	ed ting ribing	Kocader	Discoloration resistance	Light	Decolarizing property
	RE-1 ((100)	DY-1	(2)	SB-1	3	WA-1	(3)	7-04	(2)	9	(8)	2			
7	_	(001)	DY-1		SB-2	3	WA-1	(2)	S	(0.6)	V		ä) נ	2 6	< -
6	-	100	DY-2	(2)		(9)	WA-	e	A0-7	(3)	AD-5		2) <i>c</i>	a (٠.
•	RE-6	(S	DY-2	(3)		e	WA-	9	9 0 V	3	9	9	ă	: U) #	٠.
9	•	100)	DY-1		SR-1	(3)	WA-1	ε	11-04	3	Ą		3	Ų	1 4	: <
•	-	(100)	DY-1			9	WA-1	(5)	A0-6	3	AD-6		X	0	1 (< <
7	-	8	DY-3	3	SE-1	3	I-VA	9	1		VD 6		ğ	· <) <	: 4
89	_	100	DY-3	8		€	MA-I	3	1		AD-5		ğ	! «	: •	ء د
G	_	(100)	<u>0</u>	3		9	WA-1	3	1		404		ž	! <	•	ء د
10		(06)	<u>-</u> X	3		(8)	WA-1	€	•		AD-5		ă	: =	ć c	۰ د
		<u>6</u>											İ	ì	2	¢
==	RB-1	(96)	DY-4	(3)	<u>-88</u>	€	WA-1 (4)	€	ı		AD-5	(9)	Š	8	B	<
•		9	2	Ş		;	;	;					i			
:		66		9	2	3	-	Ĵ	•		Ş	9	<u>ន</u>	2	a	<
13	-	6	DY-1	(3)	1-18	3	WA-1	(3)	A0-7	(01)	Ą	(9)	×	<	•	c
~		(00)	DY-1	3	SB-2	€	WA-2	3	90V	(15)	ð		ă	: <	: <	ء د
16	_	(00)	<u>-</u>	8	SE-3	3	WA-1	3	40-11	600	4		X	! «	: <	ء د
16	RB-1	60	DY-I	(2)	SR	(4)	WA-1	(3)	ı		4		ž	: c	: ‹	٠ د

As is clear from the results shown in Tables 22 and 23, it can be understood that the loners obtained in
Experiments 1 to 20, which were prepared by using previously prepared master batches have remarkably
excellent physical properties in comparison with the loners of Comparative Experiments 1 to 16, which were
prepared by healing to melt and kneading at a time raw materials for a loner.

Also, it can be understood that when a resin binder having a polar group is used, an obtained toner shows improved excellent physical properties as well as a toner not containing a resin binder having a polar group even through a light lastness stabilizer is not used.

In addition to the ingredients used in the Examples, other ingredients can be used in the Examples as set forth in the specification to obtain substantially like same results.

Claims

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1. A decolorizable toner comprising

(A) a resin binder,

(B) at least one near infrared ray - absorbing dye selected from the group consisting of a near infrared ray - absorbing dye represented by the general formula (I);

wherein X⁻ is a halogen ion, perchloric acid ion, PF $_{\epsilon}^-$, SbF $_{\epsilon}^-$, OH⁻, sulfonic acid ion or BF $_{\epsilon}^-$, Y⁺ is a cation having absorptions in the near infrared region, and a near infrared ray - absorbing dye represented by the general formula (II):

wherein each of R¹, R², R³ and R⁴ is independently hydrogen atom, a hydrocarbon group or a hydrocarbon group containing a hetero atom, Y⁴ is the same as defined above.

(C) a decolorizing agent represented by the general formula (III):

wherein each of R⁵, R⁶, R⁷ and R⁸ is independently an alkyl group, an aryl group, an allyl group, an aralkyl group, an alkenyl group, an alkynyl group, silyl group, an alicyclic group, a substituted alkyl group, a substituted aryl group, a substituted alkyl group, a substituted aryl group, a substituted alkynyl group or a substituted silyl group, with the proviso that at least one of R⁵, R⁶, R⁷ and R⁸ is an alkyl group having 1 to 12 carbon atoms; and each of R⁵, R¹⁰, R¹¹ and R¹² is independently hydrogen atom, an alkyl group, an aryl group, an alkyl group, an alkynyl group, an alkynyl group, an alkyl group, a substituted alkyl group, a substituted aryl group, a substituted alkynyl group, and

(D) a light fastness stabilizer.

- The decolorizable toner of Claim 1, wherein said resin binder contains a resin having at least one
 functional group selected from hydroxyl group, ciano group, carboxyl group, carboxyl group and kelone
 group.
- The decolorizable toner of Claim 1, wherein said resin binder has a melt viscosity of at most 10⁷ poise
 at a temperature of 110°C and a melt viscosity of at least 10⁷ poise at a temperature of 140°C.
 - 4. The decolorizable toner of Claim I, wherein said resin binder has a light transmittance of at least 80 %.

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